

84129

Delaval, Jan

From: Roark, Jessica
Sent: Monday, January 13, 2003 3:19 PM
To: Delaval, Jan
Subject: 09/728,911

Jan,

Please search, including pending, the following from 09/728,911:

SEQ ID NO:2
SEQ ID NO:2 as an oligo
SEQ ID NO:34
SEQ ID NO:35 and
SEQ ID NO:36.

Results on paper please.

Thanks!

Jessica H. Roark

CM1 8A03
Mailbox 9E12
Art Unit 1644
703 605-1209

Jan Delaval
Reference Librarian
Biotechnology & Chemical Library
CM1 1E07 - 703-308-4498
jan.delaval@uspto.gov



f . 2

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: January 13, 2003, 15:34:21 ; Search time 35 seconds
(without alignments)
879.454 Million cell updates/sec

Title: US-09-728-911-2

Perfect score: 231

Sequence: 1 MPRHCFGLISFLTGVA.....YQPLDRSQSRSEECVEIP 231

Scoring table: OLIGO
Gapop 60.0, Gapext 60.0

Searched: 908470 seqs, 133250620 residues

Wc size: 0

Total number of hits satisfying chosen parameters: 908470

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 100 summaries

Database: A_Geneseq_101002.*

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3: /SID2/gcgdata/geneseq/geneseq-emb1/AA1982.DAT:*
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22: /SID2/gcgdata/geneseq/geneseq-emb1/AA2001.DAT:*
23: /SID2/gcgdata/geneseq/geneseq-emb1/AA2002.DAT:*
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	231	100.0	231	22	AAE05048
2	231	100.0	231	22	AAE02460
3	231	100.0	231	22	AAE02460
4	231	100.0	231	23	AAO17381
5	231	100.0	231	23	AAU00003
6	231	100.0	231	23	ABG34086
7	231	100.0	231	23	AAE17320
8	212	91.8	214	23	AAE17319
9	210	90.9	210	22	AAE02460
10	165	71.4	262	22	AAU09186

11	165	71.4	263	23	AAU080324	Human IL-TIF/IL-22
12	165	71.4	263	23	AAE17321	Human cytokine rec
13	154	66.7	263	23	AAO17382	Human cytokine rec
14	105	45.5	249	22	AAE02458	Human DNAX cytokin
15	105	45.5	249	22	AAO17380	Human DNAX cytokin
16	97	42.0	130	22	AAE02461	Human cytokine rec
17	56	24.2	56	22	ABE36621	Human DNAX cytokin
18	56	24.2	56	22	ABE36621	Human DNAX cytokin
19	56	24.2	56	22	ABE36621	Human DNAX cytokin
20	56	24.2	56	22	ABE36621	Human DNAX cytokin
21	56	24.2	56	22	ABE36621	Human DNAX cytokin
22	56	24.2	56	22	ABE36621	Human DNAX cytokin
23	56	24.2	56	22	ABE36621	Human DNAX cytokin
24	56	24.2	56	22	ABE36621	Human DNAX cytokin
25	56	24.2	56	22	ABE36621	Human DNAX cytokin
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28	56	24.2	56	22	ABE36621	Human DNAX cytokin
29	56	24.2	56	22	ABE36621	Human DNAX cytokin
30	56	24.2	56	22	ABE36621	Human DNAX cytokin
31	56	24.2	56	22	ABE36621	Human DNAX cytokin
32	56	24.2	56	22	ABE36621	Human DNAX cytokin
33	56	24.2	56	22	ABE36621	Human DNAX cytokin
34	56	24.2	56	22	ABE36621	Human DNAX cytokin
35	56	24.2	56	22	ABE36621	Human DNAX cytokin
36	56	24.2	56	22	ABE36621	Human DNAX cytokin
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39	56	24.2	56	22	ABE36621	Human DNAX cytokin
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44	56	24.2	56	22	ABE36621	Human DNAX cytokin
45	56	24.2	56	22	ABE36621	Human DNAX cytokin
46	56	24.2	56	22	ABE36621	Human DNAX cytokin
47	56	24.2	56	22	ABE36621	Human DNAX cytokin
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51	56	24.2	56	22	ABE36621	Human DNAX cytokin
52	56	24.2	56	22	ABE36621	Human DNAX cytokin
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54	56	24.2	56	22	ABE36621	Human DNAX cytokin
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60	56	24.2	56	22	ABE36621	Human DNAX cytokin
61	56	24.2	56	22	ABE36621	Human DNAX cytokin
62	56	24.2	56	22	ABE36621	Human DNAX cytokin
63	56	24.2	56	22	ABE36621	Human DNAX cytokin
64	56	24.2	56	22	ABE36621	Human DNAX cytokin
65	56	24.2	56	22	ABE36621	Human DNAX cytokin
66	56	24.2	56	22	ABE36621	Human DNAX cytokin
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84 6 2.6 58 22 AAMB9102 Human immune/haena
 85 6 2.6 59 21 AAY76330 Fragment of human
 86 6 2.6 59 22 AAO09410 Human polypeptide
 87 6 2.6 61 23 ABP03236 Human ORFX protein
 88 6 2.6 64 22 AAU52005 Propionibacterium
 89 6 2.6 64 22 AAG74813 Human colon cancer
 90 6 2.6 65 22 ABB96568 Human testicular a
 91 6 2.6 65 22 AAM96564 Human reproductive
 92 6 2.6 66 21 AAG41178 Zea mays protein f
 93 6 2.6 66 22 AAU46335 Propionibacterium
 94 6 2.6 67 22 AAO13356 Human polypeptide
 95 6 2.6 68 22 AAU20237 Human novel endocr
 96 6 2.6 70 22 AAM92880 Human digestive sy
 97 6 2.6 73 22 ABG18265 Novel human diagno
 98 6 2.6 76 19 AAY20295 Human apolipoprote
 99 6 2.6 76 22 AAM82738 Human immune/haena
 100 6 2.6 77 22 ABG26679 Novel human diagno

ALIGNMENTS

RESULT 1
 AAE05048
 ID AAE05048 standard; Protein; 231 AA.
 XX
 AC AAE05048;
 XX

DT 10-SEP-2001 (first entry)
 XX

DE Human ZCYTO18 soluble receptor antagonist, zcytor16 protein.
 XX

KW Human; cytostatic; cytokine; ZCYTO18 protein; genetic abnormality;
 KW cancer; inflammation; gene therapy; zcytor16.
 XX

OS Homo sapiens.
 XX

PN WO200146422-A1.
 XX

PD 28-JUN-2001.
 XX

PP 22-DEC-2000; 2000WO-US35308.
 XX

PR 23-DEC-1999; 99US-0471767.
 PR 01-DEC-2000; 2000US-0250841.
 XX

PA (ZYMO) ZYMOGENETICS INC.
 XX

PI Presnell SR, Kindsvogel W;
 XX

DR WPI; 2001-408648/43.
 DR N-PSDB; AAD09745.
 XX

PT Novel human cytokine polypeptide, ZCYTO18, useful for treating cancer -
 XX

PS Example 13A; Page 158-159; 167pp; English.
 XX

CC The patent discloses novel human cytokine, ZCYTO18 protein and its
 CC corresponding DNA. ZCYTO18 protein induces proliferation of cells
 CC expressing zcytor1, a receptor for ZCYTO18 or induces cytotoxicity
 CC in K562 cells. ZCYTO18 DNA is useful for detecting a genetic
 CC abnormality in a patient. ZCYTO18 DNA and its antibodies are useful
 CC for detecting cancer and inflammation. ZCYTO18 protein is useful for
 CC killing cancer cells. It is useful for increasing platelets in a
 CC patient or injured tissue. It is also used in gene therapy.
 CC The present sequence is human zcytor16, which is a naturally expressed
 CC soluble receptor antagonist of ZCYTO18 protein.
 XX

SQ Sequence 231 AA;
 XX

Query Match 100.0%; Score 231; DB 22; Length 231;
 Best Local Similarity 100.0%; Pred. No. 9.7e-220;
 Matches 231; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MPMKHCFLGFLISFFLTGVAGTQSTHESLKPORVQFSRNFHNILOWQFGRALTGNSSVY 60
 DB 1 MPMKHCFLGFLISFFLTGVAGTQSTHESLKPORVQFSRNFHNILOWQFGRALTGNSSVY 60
 QY 61 FVOYKIYGORQWKNKEDCWGTQELSCDLTSETSDIOEPYIGRVRAASAGSYSEWSMTRF 120
 DB 61 FVOYKIYGORQWKNKEDCWGTQELSCDLTSETSDIOEPYIGRVRAASAGSYSEWSMTRF 120
 QY 121 TPWWTETKIDPPWNMTQVNGSLVILHAPNLPYQKEKNVSIEDYVYLLVRFVFINNSL 180
 DB 121 TPWWTETKIDPPWNMTQVNGSLVILHAPNLPYQKEKNVSIEDYVYLLVRFVFINNSL 180
 QY 181 EKEQKYVEGAHRAVEIALTPHSSYCVVAEIIYQPMLEDRRSORSEERCVEIP 231
 DB 181 EKEQKYVEGAHRAVEIALTPHSSYCVVAEIIYQPMLEDRRSORSEERCVEIP 231

RESULT 2

AAE02460
 ID AAE02460 standard; Protein; 231 AA.
 XX

AC AAE02460;
 XX

DT 10-AUG-2001 (first entry)
 XX

DE Human DNAX cytokine receptor subunit 4.2 (DCRS4.2).
 XX

KW Human; immunomodulator; DNAX cytokine receptor subunit 4.2; DCRS4.2;
 KW therapy; immunological disorder; drug screening; cell development;
 KW chromosome 6q24.1-25.2.
 XX

OS Homo sapiens.
 XX

FH Key Location/Qualifiers
 FT Peptide 1..21
 FT Protein /label= Signal-peptide
 FT 22..231
 FT /label= DCRS4.2
 FT /note= "Human mature DNAX cytokine receptor
 FT subunit 4.2"

PN WO200136467-A2.
 XX

PD 25-MAY-2001.
 XX

PP 16-NOV-2000; 2000WO-US31363.
 XX

PR 18-NOV-1999; 99US-0443060.
 PR 13-DEC-1999; 99US-0170320.
 XX

PA (SCHE) SCHERING CORP.
 XX

PI Gorman DM;
 XX

DR WPI; 2001-343800/36.
 DR N-PSDB; AAD06414.
 XX

PT New mammalian receptor proteins related to cytokine receptors, useful
 PT for regulating cell development and for diagnosis and treatment of
 PT immunological disorders -
 XX

PS Claim 3; Page 23; 124pp; English.
 XX

CC The present sequence is human DNAX cytokine receptor subunit 4.2
 CC (DCRS4.2). DCRS4 gene is located on chromosome 6q24.1-25.2.
 CC Cytokine receptors, fragments and antibodies are useful for treating
 CC immunological disorders. DCRS3 (50R), DCRS4 (cytor) or fragments are
 CC useful in drug screening to identify compounds having binding affinity
 CC to the receptor subunit. Modulators of DCRS are useful for modulating
 CC the physiology or development of a cell or tissue culture cells. A
 CC purified DCRS is useful as a reagent to detect antibodies generated in
 CC response to the presence of elevated levels of expression, or

immunological disorders which lead to production of antibody to the endogenous receptor. Cytokine receptor sequences are useful as probes for detecting levels of the cytokine receptor in patients suspected of having an immunological disorder. Antibodies have therapeutic value as useful as potent antagonists, in detecting or quantifying ligands, for isolating cDNA proteins and peptides, to screen expression libraries for particular expression products, to raise anti-idiotypic antibodies and for detecting or diagnosing various immunological conditions related to expression of the protein or cells which express the protein.

SQ **Sequence** **231 AA;**

Query Match	100.0%;	Score 231;	DB 22;	Length 231;
Best Local Similarity	100.0%;	Pred. No. 9.7e-220;		
Matches 231; Conservative	0;	Mismatches	0;	Indels 0; Gaps 0;

Qy	1	MMRHOEFLFLLSFYLTGAGTOSTHSLKRPQVQPSNRNFHLLIOWDQGRALTGNSSVY	60
Db	1	MMRHOEFLFLLSFYLTGAGTOSTHSLKRPQVQPSNRNFHLLIOWDQGRALTGNSSVY	60
Qy	61	FVQYKIVGORQKKNKEDCMGTQBLSCDITSETSDIDEPYGYRRAASAGSYSEMSMPRF	120
Db	61	FVQYKIVGORQKKNKEDCMGTQBLSCDITSETSDIDEPYGYRRAASAGSYSEMSMPRF	120
Qy	121	TPMWEKIDIPPMNITVQNGSLVITHAAPLPIRYQEKXKVSIEDYIELLRVFIINSL	180
Db	121	TPMWEKIDIPPMNITVQNGSLVITHAAPLPIRYQEKXKVSIEDYIELLRVFIINSL	180
Qy	181	EKKQKYEGAHRAVELIATLTPHSSYCVAAEYQPMIDRNSQSEERCEVPI	231
Db	181	EKKQKYEGAHRAVELIATLTPHSSYCVAAEYQPMIDRNSQSEERCEVPI	231

```

RESULT 3
AAB62657
ID AAB62657 standard; Protein; 231 AA
....

```

AC AAB62657;

DT 23-JUL-2001 (first entry)

DE Human cytokine receptor, zcytor16

KM Cytokine receptor; zcytor16; IL-T1F; antiinflammatory; cyostatic
KM antirheumatic; antiarthritic; antiasthmatic; antiatherosclerotic;
KM immunosuppressive; chromosome 6q24.1-25.2; human.

OS Homo sapiens.

FT	key	Location/Qualifiers
FT	Domain	22..108
FT		/note= "Ig domain 1"
FT	Domain	22..231
FT		/note= "extracellular domain"
FT	Domain	112..210
FT		/note= "Ig domain 2"

PN WO200140467-A1

PD 07-JUN-2001.

PF 01-DEC-2000; 2000WO-US32703.

PR 03-DEC-1999; 99US-0169049.

PR 31-OCT-2000; 2000US-0244610.

PA (ZYMO) ZYMOGENETICS INC.

PA (ZYMO) ZYMOGENETICS INC.

PI Presnell SR, Xu W, Kindsvogel W, Chen Z; ...

DR WPI; 2001-356158/37.

XX New soluble cytokine receptor polypeptides and polynucleotides, useful
PT for diagnosing and treating cancer and inflammatory conditions -
PT
XX
XX Claim 1; Page 186-188; 210pp; English.
PS
XX

The invention relates to a human cytokine receptor polypeptide, designated zcytor16. The zcytor16 polypeptide can be expressed by CC standard recombinant methodology and can bind to IL-11F (undefined). The CC zcytor16 protein is useful for: inhibiting IL-11F induced poliferation CC or differentiation of hepatoprotetic cell(s) (progenitors); reducing CC IL-11F induced or IL-9 induced inflammation, and suppressing an CC inflammatory response in a mammal with inflammation. Heteromeric/ CC multimeric receptor polypeptides such as soluble zcytor 16/CRF-4 can be CC used to reduce progression and symptoms of cancer. Zcytor16 polypeptides CC can also be used to detect IL-11F levels which is indicative of CC pathological conditions including inflammatory states (e.g. rheumatoid CC arthritis) and cancer. Antibodies that bind zcytor16 polypeptides and the CC polypeptides themselves are useful for the treatment of inflammation, CC inflammatory diseases (e.g. infection, asthma, inflammatory bowel CC disease, rheumatoid arthritis and atherosclerosis) and autoimmune CC diseases. The antibodies and zcytor16 polynucleotides are also useful CC for detecting cancer. The present sequence represents the human CC zcytor16 protein.

SQ	Sequence	231	AA;
----	----------	-----	-----

Query Match	100.0%	Score 231;	DB 22;	Length 231;
Best Local Similarity	100.0%	Pred. No. 9.7e-220;		
Matches 231; Conservative	0;	Mismatches	0;	Indels 0

Qy	1	MMRHOEIGLFLISPFELTUVACGOSTHSLKPKQVQVQSNFNHLLQMGQALGNSSVY	60
Ds	1	MPRHGCLGFLISFPLTGVAGTOSTHSLKPKQVQVQSNFNHLLQMGQALGNSSVY	60
Qy	61	FVQYKLYIGORQKMKNEDEMGTOELSCDLTSETSDIOEPYGVGRVAASAGSYSEMSMPRF	120
Ds	61	FVQYKLYIGORQKMKNEDEMGTOELSCDLTSETSDIOEPYGVGRVAASAGSYSEMSMPRF	120
Qy	121	TPMWEKTKIDPPVNNITVONGSLVYLHAPNLPYRYQKEKNSIEDYELLRYVFIINNSL	180
Ds	121	TPMWEKTKIDPPVNNITVONGSLVYLHAPNLPYRYQKEKNSIEDYELLRYVFIINNSL	180
Qy	181	EKKQKYEGAHRAVELTALTHSSYCVAAEYQPMIDRRSQSSEKCEVP 231	
Ds	181	EKKQKYEGAHRAVELTALTHSSYCVAAEYQPMIDRRSQSSEKCEVP 231	

RESULT 4
AA017381
ID AA017381 standard; Protein; 231 AA

AC AA017381

DT 08-AUG-2002 (first entry)

Human cytokine receptor variant 2

KM Human; cytokine receptor; immune disease; psoriasis; cancer; infection
KM Rheumatoid arthritis; multiple sclerosis; Crohn's disease;
KM ulcerative colitis; transplant rejection; abortion; antipsoriatic;
KM immunosuppressive; antineumatic; antiarthritic; neuroprotective;
KM antiinflammatory; anticulcer; cytostatic; dermatological;
KM chromosome 6q24.1-25.2; receptor.

OS Homo sapiens

PN EP1191035-A2

PD 27-MAR-2002

PF 24-AUG-2001; 2001EP-0250307

PR 25-SEP-2000; 2000DE-1048626.
 PR 17-NOV-2000; 2000DE-1058907.
 PR 19-DEC-2000; 2000DE-1064906.
 XX (SCHD) SCHERING AG.
 PA Weiss B, Sabat R, Assadullah K, Toshi L;
 XX
 PI
 XX
 XX
 XX
 DR WPI; 2002-332210/37.
 DR N-PSDB; AAL46000.
 XX
 XX New nucleic acid encoding soluble cytokine receptor, useful for
 PT diagnosis and treatment of e.g. immune disease, also related protein
 PT and antibodies
 XX
 XX
 PS Claim 6; Page 14; 21pp; German.
 XX
 XX The present invention provides the protein and coding sequences of 3
 CC variants of a human cytokine receptor. The sequences can be used in the
 CC diagnosis, prevention and treatment of immune diseases, including
 CC psoriasis, cancer, chronic/life-threatening infections, rheumatoid
 CC arthritis, multiple sclerosis, Crohn's disease, ulcerative colitis and
 CC transplant rejection and in reproductive medicine, e.g. for diagnosing
 CC abnormal immune reactions which cause abortions. The present sequence is
 CC variant 2 of the invention.
 XX
 SQ Sequence 231 AA;
 Query Match 100.0%; Score 231; DB 23; Length 231;
 Best Local Similarity 100.0%; Pred. No. 9.7e-220; Indels 0; Gaps 0;
 Matches 231; Conservative 0; Mismatches 0;
 QY 1 MNPKHCFLGLISFFLTGVAGTQSTHESLKPVQVQSRNFHNLQWQGRALTGNSVY 60
 DB 1 MNPKHCFLGLISFFLTGVAGTQSTHESLKPVQVQSRNFHNLQWQGRALTGNSVY 60
 QY 61 FVQYKIYGORQWKNKEDCWGTQELSCDLTSETSDIQEYVGRVRAASAGSYSEWSMTPRF 120
 DB 61 FVQYKIYGORQWKNKEDCWGTQELSCDLTSETSDIQEYVGRVRAASAGSYSEWSMTPRF 120
 QY 121 TPWWTETKIDPPVNMNITQVNGSLVLVILHAPNLPYRYQKEKNVSIEDYELLVRFVFINNSL 180
 DB 121 TPWWTETKIDPPVNMNITQVNGSLVLVILHAPNLPYRYQKEKNVSIEDYELLVRFVFINNSL 180
 QY 181 EKEQKYEGAHRAVEIEALTPHSSYCVVAEIIYQPMMLDRRSQRSEERCVEIP 231
 DB 181 EKEQKYEGAHRAVEIEALTPHSSYCVVAEIIYQPMMLDRRSQRSEERCVEIP 231
 RESULT 5
 AAU80000
 ID AAU80000 standard; Protein; 231 AA.
 XX
 AC AAU80000;
 XX
 XX 15-JUL-2002 (first entry)
 XX
 XX Human IL-TIF/IL-22 binding protein #1.
 XX
 XX Human; soluble protein; interleukin-TIF/IL-22; IL-TIF/IL-22; IL-22BP;
 KW IL-TIF/IL-22 antagonist.
 XX
 XX Homo sapiens.
 OS
 PN WO200224912-A2.
 XX
 PD 28-MAR-2002.
 XX
 XX 21-SEP-2001; 2001WO-US295976.
 PF
 XX 22-SEP-2000; 2000US-234583P.
 PR
 PR 03-NOV-2000; 2000US-245495P.
 PR 31-JUL-2001; 2001US-091916Z.

XX
 PA (LUDW-) LUDWIG INST CANCER RES.
 XX
 PI Renauld J, Dumoutier L;
 XX
 XX WPI; 2002-383190/41.
 DR N-PSDB; ABK50076.
 XX
 XX Polynucleotide and polypeptide of soluble protein which binds to
 PT interleukin-TIF/IL-22 useful for inhibiting effect of IL-TIF/IL-22 on a
 PT cell
 XX
 XX Claim 14; Page 39; 42pp; English.
 XX
 XX The present invention relates to a new polynucleotide that encodes a
 CC soluble protein which binds to interleukin (IL)-TIF/IL-22 (also referred
 CC to as IL-22BP), where the complementary sequence of the invention
 CC hybridises under stringent conditions to a nucleotide sequence of 2271
 CC or 2366 base pairs, as given in the specification. The molecules of the
 CC invention are useful for inhibiting (antagonising) effect of IL-TIF/IL-22
 CC on a cell, for determining whether IL-TIF/IL-22 is present in a sample,
 CC for inhibiting binding of IL-TIF/IL-22 to a binding partner, preferably
 CC in vitro, and for obtaining an antibody molecule specific for the soluble
 CC binding protein of the invention, from a population or panel of antibody
 CC molecules of diverse binding specificity. The soluble protein is further
 CC useful in manufacture of a medicament for treating an IL-22 mediated
 CC disease and for assaying an agent, preferably an antibody or a peptide
 CC fragment of IL-TIF/IL-22 or the soluble protein, that modulates binding
 CC of the soluble protein to IL-TIF/IL-22, where the agent identified is
 CC used in the manufacture of medicament for treating IL-TIF/IL-22 mediated
 CC disorder. The antibody is useful for determining presence of the soluble
 CC protein, where the antibody is detectably labelled. The present amino
 CC acid sequence represents the human IL-TIF/IL-22 binding protein #1 of
 CC the invention.
 XX
 SQ Sequence 231 AA;
 Query Match 100.0%; Score 231; DB 23; Length 231;
 Best Local Similarity 100.0%; Pred. No. 9.7e-220; Indels 0; Gaps 0;
 Matches 231; Conservative 0; Mismatches 0;
 QY 1 MNPKHCFLGLISFFLTGVAGTQSTHESLKPVQVQSRNFHNLQWQGRALTGNSVY 60
 DB 1 MNPKHCFLGLISFFLTGVAGTQSTHESLKPVQVQSRNFHNLQWQGRALTGNSVY 60
 QY 61 FVQYKIYGORQWKNKEDCWGTQELSCDLTSETSDIQEYVGRVRAASAGSYSEWSMTPRF 120
 DB 61 FVQYKIYGORQWKNKEDCWGTQELSCDLTSETSDIQEYVGRVRAASAGSYSEWSMTPRF 120
 QY 121 TPWWTETKIDPPVNMNITQVNGSLVLVILHAPNLPYRYQKEKNVSIEDYELLVRFVFINNSL 180
 DB 121 TPWWTETKIDPPVNMNITQVNGSLVLVILHAPNLPYRYQKEKNVSIEDYELLVRFVFINNSL 180
 QY 181 EKEQKYEGAHRAVEIEALTPHSSYCVVAEIIYQPMMLDRRSQRSEERCVEIP 231
 DB 181 EKEQKYEGAHRAVEIEALTPHSSYCVVAEIIYQPMMLDRRSQRSEERCVEIP 231
 RESULT 6
 ABG34086
 ID ABG34086 standard; Protein; 231 AA.
 XX
 AC ABG34086;
 XX
 XX 15-JUL-2002 (first entry)
 XX
 XX Human Pro peptide #57.
 DE
 XX Human; PRO; secreted protein; transmembrane protein;
 KW genetic disorder; tumour; cancer.
 XX
 OS Homo sapiens.
 XX

immunological response. The sequences of the invention are useful for treating cancers, infections, autoimmune disorders, haematopoietic disorders, wound healing disorders, cholesterol ester storage disease, inflammation, congenital muscular dystrophy, junctional epidermolysis bullosa, Parkinson's disease, Huntington's chorea, multiple sclerosis, viral and bacterial infections, Alzheimer's disease, asthma, arthritis, allergies, schizophrenia, sbg44245PROA-associated disorders, septicemia, psoriasis, inflammatory bowel disease, acute respiratory disease graft versus host disease, ischaemia, stroke, acute respiratory disease syndrome, restenosis, brain injury, AIDS, bone diseases, atherosclerosis, brain disorders including parasupranuclear palsy, myotonic dystrophy, depression, anxiety disorders and sleep disorders, cardiovascular diseases including congestive heart failure and myocardial infarction, respiratory diseases including chronic obstructive pulmonary disease, acute bronchitis and adult respiratory distress syndrome, liver disorders including hypercholesterolaemia, hypertriglyceridaemia, cirrhosis, viral and non-viral hepatitis, type II diabetes mellitus, renal disease including acute and chronic renal failure, glomerulonephritis, Fanconi's syndrome, cystinuria, skeletal muscle disorders including hypoglycaemia and tendinitis, gastrointestinal diseases including intestinal obstruction and tropical sprue, spleen disorders including hypersplenism, Hodgkin's disease and malignant lymphoma, testicular cancer, male reproductive diseases including low testosterone and male infertility. The present sequence is human cytokine receptor.

XX Sequence 231 AA;

Query Match 100.0%; Score 231; DB 23; Length 231;
 Best Local Similarity 100.0%; Pred. No. 9.7e-220;
 Matches 231; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MNPKHCFLGFLISFFLTGAGTQSTHESLKQVQFQSRNFHNLQWPGRLTGNSSVY 60
 DB 1 MNPKHCFLGFLISFFLTGAGTQSTHESLKQVQFQSRNFHNLQWPGRLTGNSSVY 60

QY 61 FVOYKIYQORQWKNKEDCWGTQELSDLTSETSDIQEPIYGRVRAASAGSSEWMTPRF 120
 DB 61 FVOYKIYQORQWKNKEDCWGTQELSDLTSETSDIQEPIYGRVRAASAGSSEWMTPRF 120

QY 121 TPWETKIDPPVNNITQVNGSLVILHAPNLPYRQKKNVSIEDYELLRVFIINSL 180
 DB 121 TPWETKIDPPVNNITQVNGSLVILHAPNLPYRQKKNVSIEDYELLRVFIINSL 180

QY 181 EXEQKYVEGAHRAVEIALTPHSSCVVAEIIYQPMLEDRSRQSERCVEIP 231
 DB 181 EXEQKYVEGAHRAVEIALTPHSSCVVAEIIYQPMLEDRSRQSERCVEIP 231

RESULT 8

AAEL7319

ID AAEI7319 standard; Protein; 214 AA.

XX AAEI7319;

XX AAEI7319;

DT 18-APR-2002 (first entry)

XX Human cytokine receptor protein, sbg456548Cytora #1.

Human; therapy; wound healing disorder; vaccine; cancer; infection; autoimmune disorder; haematopoietic disorder; inflammation; arthritis; Parkinson's disease; Huntington's chorea; schizophrenia; antiarrhythmic; multiple sclerosis; Alzheimer's disease; analgesic; cardiant; asthma; ischaemia; stroke; AIDS; bone disease; atherosclerosis; brain disorder; depression; cardiovascular disease; myocardial infarction; renal failure; respiratory disease; liver disorder; Fanconi's syndrome; spleen disorder; type II diabetes mellitus; skeletal muscle disorder; immunosuppressive; hypersplenism; renal disease; hypoglycaemia; gastrointestinal disease; neutropenic; cirrhosis; Hodgkin's disease; neuroleptic; antiinflammatory; haemostatic; vulnery; anticonvulsant; antirheumatic; neuroprotective; nephrotropic; hypotensive; vasotropic; cytostatic; cerebroprotective; allergy; cytokine receptor.

XX Homo sapiens.

OS

XX WO200198342-A1.

XX 27-DEC-2001.

XX 22-JUN-2001; 2001WO-US19929.

XX 22-JUN-2000; 2000US-213156P.

XX 22-JUN-2000; 2000US-213161P.

XX (SMIK) SMITHKLINE BEECHAM CORP.

XX (SMIK) SMITHKLINE BEECHAM PLC.

XX (GLAX) GLAXO GROUP LTD.

XX Aggarwal P, Cogswell JP, Kabnic KS, Lai Y, Martensen SA;

XX Murdock PR, Smith RF, Strum JC, Xiang Z, Xie Q, Rizni SK;

XX WPI: 2002-139783/18.

XX N-PSDB; AAD27814.

XX Novel secreted and membrane-associated polypeptides and polynucleotides

XX useful for preventing, ameliorating or correcting dysfunction or

XX disease including diabetes, cancer, hypertension and growth

XX abnormalities

XX Claim 1; Page 122; 138pp; English.

XX The invention relates to secreted and membrane-associated polypeptides

XX and polynucleotides. The sequences of the invention are useful in

XX diagnostic assays for detecting diseases associated with inappropriate

XX activity or levels of these polynucleotides, and in identifying their

XX agonists and antagonists that are potentially useful in therapy. The

XX sequences of the invention are useful as vaccines for inducing for

XX immunological response. The sequences of the invention are useful for

XX treating cancers, infections, autoimmune disorders, haematopoietic

XX disorders, wound healing disorders, cholesterol ester storage disease,

XX inflammation, congenital muscular dystrophy, junctional epidermolysis

XX bullosa, Parkinson's disease, Huntington's chorea, multiple sclerosis,

XX viral and bacterial infections, Alzheimer's disease, asthma, arthritis,

XX septicemia, psoriasis, inflammatory bowel disease, stroke, acute respiratory

XX disease, graft versus host disease, ischaemia, stroke, acute respiratory rejection,

XX syndrome, restenosis, brain injury, AIDS, bone diseases, atherosclerosis,

XX brain disorders including parasupranuclear palsy, myotonic dystrophy,

XX depression, anxiety disorders and sleep disorders, cardiovascular

XX diseases including congestive heart failure and myocardial infarction,

XX respiratory diseases including chronic obstructive pulmonary disease,

XX acute bronchitis and adult respiratory distress syndrome, liver disorders

XX including hypercholesterolaemia, hypertriglyceridaemia, cirrhosis, viral

XX and non-viral hepatitis, type II diabetes mellitus, renal disease

XX including acute and chronic renal failure, glomerulonephritis, Fanconi's

XX syndrome, cystinuria, skeletal muscle disorders including hypoglycaemia

XX and tendinitis, gastrointestinal diseases including intestinal

XX obstruction and tropical sprue, spleen disorders including hypersplenism,

XX Hodgkin's disease and malignant lymphoma, testicular cancer, male

XX reproductive diseases including low testosterone and male infertility.

XX The present sequence is human cytokine receptor.

XX SQ Sequence 214 AA;

Query Match 91.8%; Score 212; DB 23; Length 214;

Best Local Similarity 100.0%; Pred. No. 5.2e-201;

Matches 212; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 20 AGTQSTHESLKQVQFQSRNFHNLQWPGRLTGNSSVYFVQYKIYQORQWKNKEDCW 79

DB 3 AGTQSTHESLKQVQFQSRNFHNLQWPGRLTGNSSVYFVQYKIYQORQWKNKEDCW 62

QY 80 GTQELSCDLTSETSDIQEPIYGRVRAASAGSSEWMTPRFTPWETKIDPPVNNITQVN 139

DB 63 GTQELSCDLTSETSDIQEPIYGRVRAASAGSSEWMTPRFTPWETKIDPPVNNITQVN 122

QY 140 GSSLVILHAPNLPYRQKKNVSIEDYELLRVFIINSLSEKEQKYVEGAHRAVEIAL 199


```

Db 123 GLLVILHAPNLPYRYQEKVNSIEDYELLYRVFIINNSLEKEQKVEGAHRAVEIEALP 182
Qy 200 TPSSSYCVAAEITYQPMIDRRSQRSERCEIP 231
Db 183 TPSSSYCVAAEITYQPMIDRRSQRSERCEIP 214

RESULT 9
AAB62663
ID AAB62663 standard; Protein; 210 AA.
XX
AC AAB62663;
XX
DT 23-JUN-2001 (first entry)
XX
DE Human zcytor16 extracellular domain fragment (residues 22-231).
XX
KW Cytokine receptor; zcytor16; IL-TIF; antiinflammatory; cytosolic;
KW antineumatic; antiarthritic; antiasthmatic; antiatherosclerotic;
KW immunosuppressive; chromosome 6q24.1-25.2; human.
XX
OS Homo sapiens.
XX
PN MO200140467-A1.
XX
PD 07-JUN-2001.
XX
PF 01-DEC-2000; 2000WO-US32703.
XX
PR 03-DEC-1999; 99US-0169049.
PR 13-SEP-2000; 2000US-0232219.
PR 31-OCT-2000; 2000US-0244610.
XX
PA (ZYMO ) ZYMOGENETICS INC.
XX
PI Presnell SR, Xu W, Kindsvogel W, Chen Z;
XX
DR WPI; 2001-356158/37.
XX
FT New soluble cytokine receptor polypeptides and polynucleotides, useful
FT for diagnosing and treating cancer and inflammatory conditions -
XX
PS Claim 1; Page 193; 210pp; English.
XX
CC The invention relates to a human cytokine receptor polypeptide,
CC designated zcytor16. The zcytor16 polypeptide can be expressed by
CC standard recombinant methodology and can bind to IL-TIF (undefined). The
CC zcytor16 protein is useful for: inhibiting IL-TIF induced proliferation
CC of differentiation of hematopoietic cell(s) (progenitors); reducing
CC IL-TIF induced or IL-9 induced inflammation; and suppressing an
CC inflammatory response in a mammal with inflammation. Heteromeric/
CC multimeric receptor polypeptides such as soluble zcytor 16/CRF2-4 can be
CC used to reduce progression and symptoms of cancer. Zcytor16 polypeptides
CC can also be used to detect IL-TIF levels which is indicative of
CC pathological conditions including inflammatory states (e.g. rheumatoid
CC arthritis) and cancer. Antibodies that bind zcytor16 polypeptides and the
CC polypeptides themselves are useful for the treatment of inflammation,
CC inflammatory diseases (e.g. infection, asthma, inflammatory bowel
CC disease, rheumatoid arthritis and atherosclerosis) and autoimmune
CC diseases. The antibodies and zcytor16 polynucleotides are also useful
CC for detecting cancer. The present sequence represents the human zcytor16
CC extracellular domain fragment.
XX
SQ Sequence 210 AA;

```

```

Query Match 90.9%; Score 210; DB 22; Length 210;
Best Local Similarity 100.0%; Pred. No. 4,8e-199; Indels 0; Gaps 0;
Matches 210; Conservative 0; Mismatches 0;

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```

Qy 22 TOSTHSLKQRFVQFGRNFHNIQWQGRALTGNSVYFVQYKYGQRQMKKEDCWGT 81
Db 1 TOSTHSLKQRFVQFGRNFHNIQWQGRALTGNSVYFVQYKYGQRQMKKEDCWGT 60

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Qy 82 QELSCDLTSETSDIOEPYGRVRAASAGSYSEWSMTPRFTPMWETKIDPPVNNITQVNGS 141
Db 61 QELSCDLTSETSDIOEPYGRVRAASAGSYSEWSMTPRFTPMWETKIDPPVNNITQVNGS 120
Qy 142 LVLVILHAPNLPYRYQEKVNSIEDYELLYRVFIINNSLEKEQKVEGAHRAVEIEALP 201
Db 121 LVLVILHAPNLPYRYQEKVNSIEDYELLYRVFIINNSLEKEQKVEGAHRAVEIEALP 180
Qy 202 HSSYCVAAEITYQPMIDRRSQRSERCEIP 231
Db 181 HSSYCVAAEITYQPMIDRRSQRSERCEIP 210

RESULT 10
AAU09186
ID AAU09186 standard; Protein; 262 AA.
XX
AC AAU09186;
XX
DT 16-JAN-2002 (first entry)
XX
DE Human PRO19598 polypeptide.
XX
KW Human; PRO19598; clone DNA145887; immune-related disorder;
KW inflammatory disorder; infectious disorder; immunodeficiency disorder;
KW autoimmune disorder; renal disease; demyelinating disease; skin disease;
KW neoplasia; transplantation associated disease; immunosuppressive;
KW anti-inflammatory; antiasthmatic; antidiabetic.
XX
OS Homo sapiens.
XX
PN Key
XX FH Location/Qualifiers
XX FH Peptide
XX FT 1..20
XX FT /label= Signal_peptide
XX FT Modified-site
XX FT 17..22
XX FT /note= "N-myristoylation site"
XX FT 20..25
XX FT /note= "N-myristoylation site"
XX FT Protein
XX FT 21..262
XX FT /label= Mature_PRO19598_polypeptide
XX FT Modified-site
XX FT 55..58
XX FT /note= "N-glycosylation site"
XX FT Modified-site
XX FT 165..168
XX FT /note= "N-glycosylation site"
XX FT Modified-site
XX FT 170..173
XX FT /note= "N-glycosylation site"
XX FT Modified-site
XX FT 191..194
XX FT /note= "N-glycosylation site"
XX FT Modified-site
XX FT 208..211
XX FT /note= "N-glycosylation site"
XX FT Modified-site
XX FT 220..225
XX FT /note= "N-myristoylation site"
XX
PN MO200166740-A2.
XX
PD 13-SEP-2001.
XX
PF 01-MAR-2001; 2001WO-US06666.
XX
PR 03-MAR-2000; 2000US-187202P.
PR 21-MAR-2000; 2000US-191015P.
PR 30-MAY-2000; 2000WO-US14941.
PR 05-JUN-2000; 2000US-209832P.
PR 24-AUG-2000; 2000WO-US23328.
PR 01-DEC-2000; 2000WO-US32678.
XX
PA (GENTECH ) GENENTECH INC.
XX
PI Eaton DL, Fong S, Goddard A, Godowski PJ, Grimaldi CJ, Gurney AL,
PI Tumas D, Watanabe CK, Wood WI, Zhang Z;
XX
DR WPI; 2001-625876/72.

```

DR N-PSDB; AAS15368.

XX Nucleic acids encoding PRO polypeptides, useful for detecting and

PT treating immune related diseases and disorders in mammals including

PT autoimmune diseases, inflammatory diseases and asthma -

XX

PS Claim 10; Fig 18; 122pp; English.

XX

CC The present invention relates to the isolation of 9 novel human PRO

CC polypeptides and the cDNA sequences (AAS15360-AAS15368) encoding them.

CC The novel PRO polypeptides include PRO1356, PRO1268, PRO1884, PRO3444,

CC PRO3151, PRO3322, PRO9964, PRO10008 and PRO19598. The cDNA sequences

CC encoding these PRO polypeptides have been designated as clones

CC DNA64886-1601, DNA64903-1553, DNA84318-2520, DNA87997, DNA89273,

CC DNA92223-2567, DNA96973, DNA101921 and DNA145887 respectively.

CC Compositions (e.g. vaccines) containing PRO polypeptides and methods of

CC using these compositions are useful in the treatment and diagnosis of

CC immune-related disorders. Such disorders include immune-mediated

CC inflammatory disorders (e.g. osteoarthritis), non-immune-mediated

CC inflammatory disorders (e.g. diabetes mellitus), infectious disorders

CC (e.g. granulomatous hepatitis), immunodeficiency disorders (e.g. AIDS),

CC autoimmune disorders (e.g. rheumatoid arthritis), immune-related renal

CC diseases (e.g. cirrhosis), demyelinating diseases of the peripheral or

CC central nervous system (e.g. Guillain-Barre syndrome), immune-mediated

CC skin diseases (e.g. contact dermatitis), neoplasias and transplantation

CC associated diseases. The polynucleotide sequences of the invention may

CC be used in gene therapy. AAU09178-AAU09186 represent the novel human

CC PRO polypeptides of the invention.

XX

SQ Sequence 262 AA;

Query Match 71.4%; Score 165; DB 22; Length 262;

Best Local Similarity 100.0%; Pred. No. 1.5e-154;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 67 YGQROWKNEKDCWGTQELSCDLTSETSDIQEPPYGRVRAASAGSYSEWSMTFRFTPWWT 126

Db 98 YGQROWKNEKDCWGTQELSCDLTSETSDIQEPPYGRVRAASAGSYSEWSMTFRFTPWWT 157

QY 127 KIDPPVMNITQVNGSLVLHAPNLPYRQKKNVSIEDYELLRVFVFINNSLEKEQV 186

Db 158 KIDPPVMNITQVNGSLVLHAPNLPYRQKKNVSIEDYELLRVFVFINNSLEKEQV 217

QY 187 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQRSEERCVEIP 231

Db 218 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQRSEERCVEIP 262

RESULT 11

AAU80324

ID AAU80324 standard; Protein; 263 AA.

AC AAU80324;

XX

XX 15-JUL-2002 (first entry)

DE

XX Human IL-TIF/IL-22 binding protein #2.

XX

XX Human; soluble protein; interleukin-TIF/IL-22; IL-TIF/IL-22; IL-22BP;

KW IL-TIF/IL-22 antagonist.

XX

OS Homo sapiens.

XX

XX WO200224912-A2.

PN

XX

XX 28-MAR-2002.

PD

XX

XX 21-SEP-2001; 2001WO-US29576.

XX

XX 22-SEP-2000; 2000US-234583P.

PR

XX 03-NOV-2000; 2000US-245495P.

PR

XX 31-JUL-2001; 2001US-0919162.

XX

PA (LUDW-) LUDWIG INST CANCER RES.

XX

PI Renauld J, Dumoutier L;

XX

DR WPI: 2002-383190/41.

DR N-PSDB; ABK50080.

XX

XX Polynucleotide and polypeptide of soluble protein which binds to

PT interleukin-TIF/IL-22 useful for inhibiting effect of IL-TIF/IL-22 on a

PT cell -

XX

XX Claim 14; Page 41-42; 42pp; English.

PS

CC The present invention relates to a new polynucleotide that encodes a

CC soluble protein which binds to interleukin (IL)-TIF/IL-22 (also referred

CC to as IL-22BP), where the complementary sequence of the invention

CC hybridises under stringent conditions to a nucleotide sequence of 2271

CC or 2366 base pairs, as given in the specification. The molecules of the

CC invention are useful for inhibiting (antagonising) effect of IL-TIF/IL-22

CC on a cell, for determining whether IL-TIF/IL-22 is present in a sample,

CC for inhibiting binding of IL-TIF/IL-22 to a binding partner, preferably

CC in vitro, and for obtaining an antibody molecule specific for the soluble

CC binding protein of the invention, from a population or panel of antibody

CC molecules of diverse binding specificity. The soluble protein is further

CC useful in manufacture of a medicament for treating an IL-22 mediated

CC disease and for assaying an agent, preferably an antibody or a peptide

CC fragment of IL-TIF/IL-22 or the soluble protein, that modulates binding

CC of the soluble protein to IL-TIF/IL-22, where the agent identified is

CC used in the manufacture of medicament for treating IL-TIF/IL-22 mediated

CC disorder. The antibody is useful for determining presence of the soluble

CC protein, where the antibody is detectably labelled. The present amino

CC acid sequence represents the human IL-TIF/IL-22 binding protein #2 of

CC the invention.

XX

SQ Sequence 263 AA;

Query Match 71.4%; Score 165; DB 23; Length 263;

Best Local Similarity 100.0%; Pred. No. 1.5e-154;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 67 YGQROWKNEKDCWGTQELSCDLTSETSDIQEPPYGRVRAASAGSYSEWSMTFRFTPWWT 126

Db 99 YGQROWKNEKDCWGTQELSCDLTSETSDIQEPPYGRVRAASAGSYSEWSMTFRFTPWWT 158

QY 127 KIDPPVMNITQVNGSLVLHAPNLPYRQKKNVSIEDYELLRVFVFINNSLEKEQV 186

Db 159 KIDPPVMNITQVNGSLVLHAPNLPYRQKKNVSIEDYELLRVFVFINNSLEKEQV 218

QY 187 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQRSEERCVEIP 231

Db 219 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQRSEERCVEIP 263

RESULT 12

AAE17321

ID AAE17321 standard; Protein; 263 AA.

XX

AC AAE17321;

XX

XX 18-APR-2002 (first entry)

DT

XX

XX Human cytokine receptor protein, sbg456548CytoRa #3.

DE

XX

XX Human; therapy; wound healing disorder; vaccine; cancer; infection;

KW autoimmune disorder; haematopoietic disorder; inflammation; arthritis;

KW Parkinson's disease; Huntington's chorea; schizophrenia; antarthritic;

KW multiple sclerosis; Alzheimer's disease; analgesic; cardiac; asthma;

KW ischaemia; stroke; AIDS; bone disease; atherosclerosis; brain disorder;

KW depression; cardiovascular disease; myocardial infarction; renal failure;

KW respiratory disease; liver disorder; Fanconi's syndrome; spleen disorder;

KW type II diabetes mellitus; skeletal muscle disorder; immunosuppressive;

KW hyperplenism; renal disease; hypoglycaemia; gastrointestinal disease;

KW neutropenic; cirrhosis; Hodgkin's disease; neuroleptic; antiinflammatory;

KM haemostatic; vulnerary; anticonvulsant; antirheumatic; neuroprotective;
 KM nephrotropic; hypotensive; vasotrophic; cyostatic; cerebroprotective;
 KM allergy; cytokine receptor.
 OS Homo sapiens.
 XX WO200198342-A1.
 XX
 XX PD 27-DEC-2001.
 XX
 XX PF 22-JUN-2001; 2001WO-US19929.
 XX
 XX PR 22-JUN-2000; 2000US-213156P.
 XX PR 22-JUN-2000; 2000US-213161P.
 XX
 XX PA (SMIK) SMITHKLINE BEECHAM CORP.
 XX PA (SMIK) SMITHKLINE BEECHAM PLC.
 XX PA (GLAX) GLAXO GROUP LTD.
 XX
 PI Agarwal P, Cogswell JP, Kabnic KS, Lai Y, Martensen SA;
 PI Murdoch PR, Smith RF, Strum JC, Xiang Z, Xie Q, Rizni SK;
 DR WPI: 2002-139783/18.
 DR N-PSDB; AAD27816.
 XX
 XX PS Claim 1; Page 133-134; 138pp; English.
 XX
 CC The invention relates to secreted and membrane-associated polypeptides
 CC and polynucleotides. The sequences of the invention are useful in
 CC diagnostic assays for detecting diseases associated with inappropriate
 CC activity or levels of these polynucleotides, and in identifying their
 CC agonists and antagonists that are potentially useful in therapy. The
 CC sequences of the invention are useful as vaccines for inducing
 CC immunological response. The sequences of the invention are useful for
 CC treating cancers, infections, autoimmune disorders, haematopoietic
 CC disorders, wound healing disorders, cholesterol ester storage disease,
 CC inflammation, congenital muscular dystrophy, junctional epidermolysis,
 CC bullous, Parkinson's disease, Huntington's chorea, multiple sclerosis,
 CC viral and bacterial infections, Alzheimer's disease, asthma, arthritis,
 CC allergies, schizophrenia, b9442445PROA-associated disorders,
 CC septiccaemia, psoriasis, inflammatory bowel disease, transplant rejection,
 CC graft versus host disease, ischaemia, stroke, acute respiratory disease,
 CC syndrome, restenosis, brain injury, AIDS, bone diseases, atherosclerosis,
 CC brain disorders including parasupranuclear palsy, myotonic dystrophy,
 CC depression, anxiety disorders and sleep disorders, cardiovascular
 CC diseases including congestive heart failure and myocardial infarction,
 CC respiratory diseases including chronic obstructive pulmonary disease,
 CC acute bronchitis and adult respiratory distress syndrome, liver disorders
 CC including hypercholesterolaemia, hypertriglyceridaemia, cirrhosis, viral
 CC and non-viral hepatitis, type II diabetes mellitus, renal disease
 CC including acute and chronic renal failure, glomerulonephritis, Fanconi's
 CC syndrome, cystinuria, skeletal muscle disorders including hypoglycaemia
 CC and tendinitis, gastrointestinal diseases including intestinal
 CC obstruction and tropical sprue, spleen disorders including hyperplenism,
 CC Hodgkin's disease and malignant lymphoma, testicular cancer, male
 CC reproductive diseases including low testosterone and male infertility.
 CC The present sequence is human cytokine receptor.
 CC
 XX
 SQ Sequence 263 AA;

Query Match 71.4%; Score 165; DB 23; Length 263;
 Best Local Similarity 100.0%; Pred. No. 1.5e-154;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 67 YGQORWKKEKEDCMTQELSCDLTSETSDIOEPYGRVRAAASAGSYSEWMTPTPTTMMET 126
 DB 99 YGQORWKKEKEDCMTQELSCDLTSETSDIOEPYGRVRAAASAGSYSEWMTPTPTTMMET 158

QY 127 KIDPVMNITQVNGSLVILHAPLPRYQKEKNVSIEDYELLVRFIINNSLEKEQKV 186
 DB 159 KIDPVMNITQVNGSLVILHAPLPRYQKEKNVSIEDYELLVRFIINNSLEKEQKV 218
 QY 187 YEGARAVEIALTPHSSYCVVAEIQPMLDRRSQSEECVEIP 231
 DB 219 YEGARAVEIALTPHSSYCVVAEIQPMLDRRSQSEECVEIP 263

RESULT 13

AA017382
ID AA017382 standard; Protein; 263 AA.

XX AA017382;

DT 08-AUG-2002 (first entry)

DE Human cytokine receptor variant 3.

XX Human; cytokine receptor; immune disease; psoriasis; cancer; infection;
 KM rheumatoid arthritis; multiple sclerosis; Crohn's disease;
 KM ulcerative colitis; transplant rejection; abortion; antipsoriatic;
 KM immunosuppressive; antirheumatic; antiarthritis; neuroprotective;
 KM antiinflammatory; antitumor; cyostatic; dermatological;
 KM chromosome 6q24.1-25.2; receptor.
 XX
 XX OS Homo sapiens.
 XX
 XX PN EP1191035-A2.
 XX PD 27-MAR-2002.
 XX
 XX PF 24-AUG-2001; 2001EP-0250307.
 XX
 XX PR 25-SEP-2000; 2000DE-1048626.
 XX PR 17-NOV-2000; 2000DE-1058907.
 XX PR 19-DEC-2000; 2000DE-1064906.
 XX
 XX PA (SCHD) SCHERING AG.
 XX
 XX PI Weiss B, Sabat R, Aesadullah K, Toishi L;
 XX DR WPI: 2002-332210/37.
 XX DR N-PSDB; AAL46001.
 XX
 XX PS New nucleic acid encoding soluble cytokine receptor, useful for
 XX PT diagnosis and treatment of e.g. immune disease, also related protein
 XX and antibodies -
 XX
 XX
 XX Claim 6; Page 15; 21pp; German.

CC The present invention provides the protein and coding sequences of 3
 CC variants of a human cytokine receptor. The sequences can be used in the
 CC diagnosis, prevention and treatment of immune diseases, including
 CC psoriasis, cancer, chronic/life-threatening infections, rheumatoid
 CC arthritis, multiple sclerosis, Crohn's disease, ulcerative colitis and
 CC transplant rejection and in reproductive medicine, e.g. for diagnosing
 CC abnormal immune reactions which cause abortions. The present sequence is
 CC variant 3 of the invention.
 CC
 XX
 SQ Sequence 263 AA;

Query Match 66.7%; Score 154; DB 23; Length 263;
 Best Local Similarity 100.0%; Pred. No. 1.1e-143;
 Matches 154; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 67 YGQORWKKEKEDCMTQELSCDLTSETSDIOEPYGRVRAAASAGSYSEWMTPTPTTMMET 126
 DB 99 YGQORWKKEKEDCMTQELSCDLTSETSDIOEPYGRVRAAASAGSYSEWMTPTPTTMMET 158
 QY 127 KIDPVMNITQVNGSLVILHAPLPRYQKEKNVSIEDYELLVRFIINNSLEKEQKV 186
 DB 159 KIDPVMNITQVNGSLVILHAPLPRYQKEKNVSIEDYELLVRFIINNSLEKEQKV 218

QY 187 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRS 220
 Db 219 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRS 252

RESULT 14
 AAE02458
 ID AAE02458 standard; Protein; 249 AA.
 AC AAE02458;
 XX 10-AUG-2001 (first entry)
 DT Human DNAX cytokine receptor subunit 4.1 (DCRS4.1).
 XX Human; immunomodulator; DNAX cytokine receptor subunit 4.1; DCRS4.1;
 KW therapy; immunological disorder; drug screening; cell development;
 KW chromosome 6q24.1-25.2.
 XX Homo sapiens.

OS Key Location/Qualifiers
 FH Peptide 1..21
 FT /label= Signal-peptide
 FT 22..249
 FT /label= DCRS4.1
 FT /note= "Human mature DNAX cytokine receptor
 subunit 4.1"
 FT 24
 FT /note= "CK2 phosphorylation site"
 FT 25
 FT /note= "Calcium phosphorylation site"
 FT 28
 FT /note= "PKC phosphorylation site"
 FT 31..70
 FT /label= Cytokine_receptor_domain
 FT 51
 FT /note= "cAMP PK site"
 FT 56
 FT /note= "N-glycosylated"
 FT 78..86
 FT /label= Conserved_disulphide_linkage
 FT 81
 FT /note= "Calcium phosphorylation site"
 FT 85
 FT /note= "Calcium phosphorylation site"
 FT 89
 FT /note= "Calcium phosphorylation site"
 FT 92
 FT /note= "Calcium phosphorylation site"
 FT 100
 FT /note= "Amidation site"
 FT 110
 FT /note= "Myristoyl site"
 FT 118
 FT /note= "PKC phosphorylation site"
 FT 119
 FT /note= "cAMP phosphorylation site"
 FT 119
 FT /note= "cAMP PK site"
 FT 124
 FT /note= "Myristoyl site"
 FT 127
 FT /note= "cAMP PK site"
 FT 152
 FT /note= "N-glycosylated"
 FT 157
 FT /note= "N-glycosylated"
 FT 177
 FT /note= "cAMP PK site"
 FT 178
 FT /note= "N-glycosylated"

FT Modified-site 180 /note= "Calcium phosphorylation site"
 FT Modified-site 180 /note= "CK2 phosphorylation site"
 FT Modified-site 195 /note= "N-glycosylated"
 FT Modified-site 197 /note= "Calcium phosphorylation site"
 FT Modified-site 207 /note= "Myristoyl site"
 FT Modified-site 238 /note= "PKC phosphorylation site"
 FT Modified-site 241 /note= "Calcium phosphorylation site"
 XX WO200136467-A2.
 PN 25-MAY-2001.
 XX 16-NOV-2000; 2000WO-US31363.
 XX 18-NOV-1999; 99US-0443060.
 PR 13-DEC-1999; 99US-0170320.
 XX (SCHE) SCHERING CORP.
 PA Gorman DM;
 PI WPI; 2001-343800/36.
 XX N-PSDB; AAD06410.
 DR New mammalian receptor proteins related to cytokine receptors, useful
 DR for regulating cell development and for diagnosis and treatment of
 DR immunological disorders
 PT Claim 3; Page 22; 124pp; English.
 PS The present sequence is human DNAX cytokine receptor subunit 4.1
 XX (DCRS4.1). DCRS4 gene is located on chromosome 6q24.1-25.2.
 CC Cytokine receptors, fragments and antibodies are useful for treating
 CC immunological disorders. DCRS3 (50R), DCRS4 (cytor) or fragments are
 CC useful in drug screening to identify compounds having binding affinity
 CC to the receptor subunit. Modulators of DCRS are useful for modulating
 CC the physiology or development of a cell or tissue culture cells. A
 CC purified DCRS is useful as a reagent to detect antibodies generated in
 CC response to the presence of elevated levels of expression, or
 CC immunological disorders which lead to production of antibody to the
 CC endogenous receptor. Cytokine receptor sequences are useful as probes
 CC for detecting levels of the cytokine receptor in patients suspected of
 CC having an immunological disorder. Antibodies have therapeutic value, are
 CC useful as potent antagonist, in detecting or quantifying ligands, for
 CC isolating DCRS proteins and peptides, to screen expression libraries for
 CC particular expression products, to raise anti-idiotypic antibodies and
 CC for detecting or diagnosing various immunological conditions related to
 CC expression of the protein or cells which express the protein.

SQ Sequence 249 AA;

Query Match 45.5%; Score 105; DB 22; Length 249;
 Best Local Similarity 100.0%; Pred. No. 2.3e-95;
 Matches 105; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 127 KIDPPVMNITQVNGSLVILHAPNLPYRYQKKNVSIEDYVELLYRVFVFNNSLEKEQKV 186
 Db 145 KIDPPVMNITQVNGSLVILHAPNLPYRYQKKNVSIEDYVELLYRVFVFNNSLEKEQKV 204

QY 187 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQSRSEKRCVEIP 231

Db 205 YEGAHRAVEIEALTPHSSYCVVAEIQPMLDRRSQSRSEKRCVEIP 249

RESULT 15
 AAO17380

ID AA017380.standard; Protein; 249 AA.
 AC AA017380;
 DT 08-AUG-2002 (first entry)
 XX
 DE Human cytokine receptor variant 1.
 KW Human; cytokine receptor; immune disease; psoriasis; cancer; infection;
 KW rheumatoid arthritis; multiple sclerosis; Crohn's disease;
 KW ulcerative colitis; transplant rejection; abortion; antipsoriatic;
 KW immunosuppressive; antirheumatic; antiarthritic; neuroprotective;
 KW antitumour; anticancer; cytostatic; dermatological;
 KW chromosome 6q24.1-25.2; receptor.
 XX
 OS Homo sapiens.
 PN EPI191035-A2.
 PD 27-MAR-2002.
 XX
 P1 24-AUG-2001; 2001EP-0250307.
 PR 25-SEP-2000; 2000DE-1048626.
 PR 17-NOV-2000; 2000DE-1058907.
 PR 19-DEC-2000; 2000DE-1064906.
 XX
 PA (SCHD) SCHERING AG.
 PI Weies B, Sabat R, Assadullah K, Toshi L;
 DR WPI; 2002-332210/37.
 DR N-PSDB; AAL45999.
 XX
 PT New nucleic acid encoding soluble cytokine receptor, useful for
 PT diagnosis and treatment of e.g. immune disease, also related protein
 PT and antibodies -
 XX
 PS Claim 6; Page 12-13; 21pp; German.
 CC The present invention provides the protein and coding sequences of 3
 CC variants of a human cytokine receptor. The sequences can be used in the
 CC diagnosis, prevention and treatment of immune diseases, including
 CC psoriasis, cancer, chronic/life-threatening infections, rheumatoid
 CC arthritis, multiple sclerosis, Crohn's disease, ulcerative colitis and
 CC transplant rejection and in reproductive medicine, e.g. for diagnosing
 CC abnormal immune reactions which cause abortions. The present sequence is
 CC variant 1 of the invention.
 XX
 SQ Sequence 249 AA;
 Query Match 45.5%; Score 105; DB 23; Length 249;
 Best Local Similarity 100.0%; Pred. No. 2.3e-95;
 Matches 105; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 127 KIDPPYMNITQVNGSLVILHAHNLPRYQKKNVSIIDYELLYRVFIINNSLEKQKV 186
 DB 145 KIDPPYMNITQVNGSLVILHAHNLPRYQKKNVSIIDYELLYRVFIINNSLEKQKV 204
 QY 187 YEGARHAEVEIALTPHSSYCVAAETIYOMLDRRSQSRERCEVP 231
 DB 205 YEGARHAEVEIALTPHSSYCVAAETIYOMLDRRSQSRERCEVP 249
 RESULT 16
 ID AAE02461
 AC AAE02461; standard; Protein; 130 AA.
 XX
 AC AAE02461;
 DT 10-AUG-2001 (first entry)
 XX
 DE Human DNAX cytokine receptor subunit 4.3 (DCRS4.3).

XX Human; immunomodulator; DNAX cytokine receptor subunit 4.3; DCRS4.3;
 KW therapy; immunological disorder; drug screening; cell development;
 KW chromosome 6q24.1-25.2.
 XX
 OS Homo sapiens.
 PN Key Location/Qualifiers
 FT Peptide 1..21
 FT Protein /label= Signal-peptide
 FT Protein 22..130
 FT /label= DCRS4.3
 FT /note= "human mature DNAX cytokine receptor
 subunit 4.3"
 XX
 XX WO200136467-A2.
 XX
 XX 25-MAY-2001.
 XX
 PD 16-NOV-2000; 2000WO-US31363.
 XX
 PF 18-NOV-1999; 99US-0443060.
 PR 13-DEC-1999; 99US-0170320.
 XX
 PA (SCHE) SCHERING CORP.
 XX
 PI Gorman DM;
 DR WPI; 2001-343800/36.
 DR N-PSDB; AAD06416.
 XX
 PT New mammalian receptor proteins related to cytokine receptors, useful
 PT for regulating cell development and for diagnosis and treatment of
 PT immunological disorders -
 XX
 PS Claim 3; Page 24; 124pp; English.
 CC The present sequence is human DNAX cytokine receptor subunit 4.3
 CC (DCRS4.3). DCRS4 gene is located on chromosome 6q24.1-25.2.
 CC Cytokine receptors, fragments and antibodies are useful for treating
 CC immunological disorders. DCRS3 (50R), DCRS4 (cytor) or fragments are
 CC useful in drug screening to identify compounds having binding affinity
 CC to the receptor subunit. Modulators of DCRS are useful for modulating
 CC the physiology or development of a cell or tissue culture cells. A
 CC purified DCRS is useful as a reagent to detect antibodies generated in
 CC response to the presence of elevated levels of expression, or
 CC immunological disorders which lead to production of antibody to the
 CC endogenous receptor. Cytokine receptor sequences are useful as probes
 CC for detecting levels of the cytokine receptor in patients suspected of
 CC having an immunological disorder. Antibodies have therapeutic value, are
 CC useful as potent antagonist, in detecting or quantifying ligands, for
 CC isolating DCRS proteins and peptides, to screen expression libraries for
 CC particular expression products, to raise anti-idiotypic antibodies for
 CC for detecting or diagnosing various immunological conditions related to
 CC expression of the protein or cells which express the protein.
 XX
 SQ Sequence 130 AA;
 Query Match 42.0%; Score 97; DB 22; Length 130;
 Best Local Similarity 100.0%; Pred. No. 1.1e-87;
 Matches 97; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 MNPKHCFLGFLISFLTGACTOSTHSLKPRVQFSRNFHNILOQOPRALTGNSVY 60
 DB 1 MNPKHCFLGFLISFLTGACTOSTHSLKPRVQFSRNFHNILOQOPRALTGNSVY 60
 QY 61 FVQYKTYGQRQWKXKEDCWGTQELSCDLTSETSDIOE 97
 DB 61 FVQYKTYGQRQWKXKEDCWGTQELSCDLTSETSDIOE 97
 RESULT 17
 ABB36621

AAAM34563	
ID	AAAM34563 standard; Protein; 56 AA.
XX	
XX	AAAM34563;
XX	
DT	17-OCT-2001 (first entry)
XX	
DE	Peptide #8600 encoded by probe for measuring placental gene expression.
XX	
DE	Probe; microarray; human; placenta; antenatal diagnosis;
KW	genetic disorder.
KW	
OS	Homo sapiens.
XX	
PN	WO200157272-A2.
XX	
PD	09-AUG-2001.
XX	
PF	30-JAN-2001; 2001WO-US000663.
XX	
PR	04-FEB-2000; 2000US-0180312.
PR	26-MAY-2000; 2000US-0207456.
PR	30-JUN-2000; 2000US-0608408.
PR	03-AUG-2000; 2000US-0632366.
PR	21-SEP-2000; 2000US-0234687.
PR	27-SEP-2000; 2000US-0236359.
PR	04-OCT-2000; 2000GB-0024263.
XX	
PA	(MOLE-) MOLECULAR DYNAMICS INC.
XX	
PI	Penn SG, Hanzel DK, Chen W, Rank DR;
XX	
PI	WPI; 2001-488897/53.
XX	
XX	Human genome-derived single exon nucleic acid probes useful for
PT	analyzing gene expression in human placenta -
XX	
PT	Claim 27; SEQ ID No 34832; 654pp; English.
PS	
PS	The present invention relates to single exon nucleic acid probes (SENPs)
CC	see AA131315-AA157546). The present sequence is a peptide encoded by on
CC	such probe. The probes are useful for producing a microarray for
CC	predicting, measuring and displaying gene expression in samples derived
CC	from human placenta. The probes are useful for antenatal diagnosis of
CC	human genetic disorders.
XX	
XX	Sequence 56 AA;
SQ	
	Query Match 24.2%; Score 56; DB 22; Length 56;
	Best Local Similarity 100.0%; Pred. No. 1.6e-47;
	Matches 56; Conservative 0; Mismatches 0; Indels 0; Gaps
QY	127 KIDPPVMITQVNGSLLVILHAPNLPYRYQKEKNVSIETYELLYRVFINNSLEK 182
	1 KIDPPVMITQVNGSLLVILHAPNLPYRYQKEKNVSIETYELLYRVFINNSLEK 56
DB	
RESULT 24	
ABG39407	
ID	ABG39407 standard; Peptide; 56 AA.
XX	
AC	ABG39407;
XX	
DT	19-AUG-2002 (first entry)
XX	
DE	Human peptide encoded by genome-derived single exon probe SEQ ID 29072
XX	
XX	Human; single exon probe; asthma; lung cancer; COPD; ILD;
KW	chronic obstructive pulmonary disease; interstitial lung disease;
KW	familial idiopathic pulmonary fibrosis; neurofibromatosis;
KW	tuberosus sclerosis; Gaucher's disease; Niemann-Pick disease;
KW	Hernansky-Pudlak syndrome; sarcoidosis; pulmonary haemoideriosis;
KW	pulmonary histiocytosis; lymphangioleiomyomatosis; Karagazer syndrome;

DT	19-AUG-2002	(first entry)	Human peptide encoded by genome-derived single exon probe SEQ ID 29072
XX			Human; single exon probe; asthma; lung cancer; COPD; ILD;
DE			chronic obstructive pulmonary disease; interstitial lung disease;
XX			familial idiopathic pulmonary fibrosis; neurofibromatosis;
KW			tuberosus sclerosis; Gaucher's disease; Niemann-Pick disease;
KW			Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemosiderosis;
KW			pulmonary histiocytosis; lymphangioleiomyomatosis; Karsagenr syndrome;

KW pulmonary, alveolar proteinosis; fibrocystic pulmonary dysplasia;
 KW primary ciliary dyskinesia; pulmonary hypertension;
 KW hyaline membrane disease.
 OS Homo sapiens.
 XX MO200186003-A2.
 XX
 XX 15-NOV-2001.
 XX
 XX 30-JAN-2001; 2001WO-US00665.
 XX
 XX 04-FEB-2000; 2000US-180312P.
 XX 26-MAY-2000; 2000US-207456P.
 XX 30-JUN-2000; 2000US-0608408.
 XX 03-AUG-2000; 2000US-0632366.
 XX 21-SEP-2000; 2000US-234687P.
 XX 27-SEP-2000; 2000US-236359P.
 XX 04-OCT-2000; 2000GB-0024263.
 XX
 XX (MOLE-) MOLECULAR DYNAMICS INC.
 XX Penn SG, Hanzel DK, Chen W, Rank DR;
 XX WPI; 2002-114183/15.
 XX
 XX Spatially-addressable set of single exon nucleic acid probes, used to
 XX measure gene expression in human lung samples -
 XX
 XX Claim 27; SEQ ID No 29072; 634bp; English.
 XX
 XX The invention relates to a spatially-addressable set of single exon
 XX nucleic acid probes for measuring gene expression in a sample derived
 XX from human lung comprising single exon nucleic acid probes having one of
 XX 12614 nucleic acid sequences mentioned in the specification, or their
 XX complements or the 12387 open reading frames derived from the 12614
 XX probes. Also included are a microarray comprising the novel set of
 XX probes; the novel set of probes which hybridize at high stringency to a
 XX nucleic acid expressed in the human lung; measuring gene expression in a
 XX sample derived from human lung, comprising (a) contacting the array with
 XX a collection of detectably labeled nucleic acids derived from human lung
 XX mRNA, and (b) measuring the label detectably bound to each probe of
 XX the array; identifying exons in a eukaryotic genome, comprising
 XX (a) algorithmically predicting at least one exon from genomic sequences
 XX of the eukaryote; and (b) detecting specific hybridization of detectably
 XX labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,
 XX having a fragment identical to the predicted exon, the probe is included
 XX in the above mentioned microarray; assigning exons to a single gene,
 XX comprising (a) identifying exons from genomic sequence by the method
 XX above and (b) measuring the expression of each of the exons in several
 XX tissues and/or cell types using hybridization to a single exon
 XX microarray having a probe with the exon, where a common pattern of
 XX expression of the exons in the tissues and/or cell types indicates that
 XX the exons should be assigned to a single gene; a peptide comprising one
 XX of 12011 sequences, mentioned in the specification, or encoded by the
 XX probes/open reading frames (ORF). The probes are used for gene
 XX expression analysis, and for identifying exons in a gene, particularly
 XX using human lung derived mRNA and for the study of lung diseases
 XX such as asthma, lung cancer, chronic obstructive pulmonary disease
 XX (COPD), interstitial lung disease (ILD), familial idiopathic pulmonary
 XX fibrosis, neurofibromatosis, tuberous sclerosis, Gaucher's disease,
 XX Niemann-Pick disease, Hermansky-Pudlak syndrome, sarcoidosis, pulmonary
 XX haemorrhagic disease, pulmonary histiocytosis, lymphangioleiomyomatosis,
 XX pulmonary alveolar proteinosis, Karagener syndrome, fibrocystic
 XX pulmonary dysplasia, primary ciliary dyskinesia, pulmonary hypertension
 XX and hyaline membrane disease. The present sequence is a peptide/protein
 XX encoded by a single exon probe of the invention.
 XX Note: The sequence data for this patent did not form part
 XX of the printed specification, but was obtained in electronic
 XX format directly from WIPO at
 XX ftp.wipo.int/pub/published_pct_sequences.
 XX
 XX Sequence 56 AA;

Query Match 24.2%; Score 56; DB 23; Length 56;
 Best Local Similarity 100.0%; Pred. No. 1.6e-47;
 Matches 56; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 127 KIDPVNMITOVNSSLVILHAPNLPYRYOKKNSIEDYELLRYVFIINSLK 182
 DB 1 KIDPVNMITOVNSSLVILHAPNLPYRYOKKNSIEDYELLRYVFIINSLK 56
 RESULT 25
 ABG44337
 ID ABG44337 standard; Peptide; 56 AA.
 AC ABG44337;
 XX 19-AUG-2002 (first entry)
 XX
 XX Human peptide encoded by genome-derived single exon probe SEQ ID 34002.
 DE
 XX Human; single exon probe; asthma; lung cancer; COPD; ILD;
 KW chronic obstructive pulmonary disease; interstitial lung disease;
 KW familial idiopathic pulmonary fibrosis; neurofibromatosis;
 KW tuberous sclerosis; Gaucher's disease; Niemann-Pick disease;
 KW Hermansky-Pudlak syndrome; sarcoidosis; pulmonary haemorrhagic
 KW disease; pulmonary histiocytosis; lymphangioleiomyomatosis; Karagener syndrome;
 KW pulmonary alveolar proteinosis; fibrocystic pulmonary dysplasia;
 KW primary ciliary dyskinesia; pulmonary hypertension;
 KW hyaline membrane disease.
 XX
 XX Homo sapiens.
 OS
 XX MO200186003-A2.
 XX
 XX 15-NOV-2001.
 XX
 XX 30-JAN-2001; 2001WO-US00665.
 XX
 XX 04-FEB-2000; 2000US-180312P.
 XX 26-MAY-2000; 2000US-207456P.
 XX 30-JUN-2000; 2000US-0608408.
 XX 03-AUG-2000; 2000US-0632366.
 XX 21-SEP-2000; 2000US-234687P.
 XX 27-SEP-2000; 2000US-236359P.
 XX 04-OCT-2000; 2000GB-0024263.
 XX
 XX (MOLE-) MOLECULAR DYNAMICS INC.
 XX Penn SG, Hanzel DK, Chen W, Rank DR;
 XX WPI; 2002-114183/15.
 XX
 XX Spatially-addressable set of single exon nucleic acid probes, used to
 XX measure gene expression in human lung samples -
 XX
 XX Claim 27; SEQ ID No 34002; 634bp; English.
 XX
 XX The invention relates to a spatially-addressable set of single exon
 XX nucleic acid probes for measuring gene expression in a sample derived
 XX from human lung comprising single exon nucleic acid probes having one of
 XX 12614 nucleic acid sequences mentioned in the specification, or their
 XX complements or the 12387 open reading frames derived from the 12614
 XX probes. Also included are a microarray comprising the novel set of
 XX probes; the novel set of probes which hybridize at high stringency to a
 XX nucleic acid expressed in the human lung; measuring gene expression in a
 XX sample derived from human lung, comprising (a) contacting the array with
 XX a collection of detectably labeled nucleic acids derived from human lung
 XX mRNA, and (b) measuring the label detectably bound to each probe of
 XX the array; identifying exons in a eukaryotic genome, comprising
 XX (a) algorithmically predicting at least one exon from genomic sequences
 XX of the eukaryote; and (b) detecting specific hybridization of detectably
 XX labeled nucleic acids from eukaryote lung mRNA, to a single exon probe,
 XX having a fragment identical to the predicted exon, the probe is included

CC in the above mentioned microarray; assigning exons to a single gene,
CC comprising (a) identifying exons from genomic sequence by the method
CC above and (b) measuring the expression of each of the exons in several
CC tissues and/or cell types using hybridisation to a single exon
CC microarrays having a probe with the exon, where a common pattern of
CC expression of the exons in the tissues and/or cell types indicates that
CC the exons should be assigned to a single gene; a peptide comprising one
CC of 12011 sequences, mentioned in the specification, or encoded by the
CC probes/open reading frames (ORF). The probes are used for gene
CC expression analysis, and for identifying exons in a gene, particularly
CC using human lung derived mRNA and for the study of lung diseases
CC such as asthma, lung cancer, chronic obstructive pulmonary disease
CC (COPD), interstitial lung disease (ILD), familial idiopathic pulmonary
CC fibrosis, neurofibromatosis, tuberous sclerosis, Gaucher's disease,
CC Niemann-Pick disease, Hermansky-Pudlak syndrome, sarcoidosis, pulmonary
CC haemangioendothelioma, pulmonary histiocytosis, lymphangioleiomyomatosis,
CC pulmonary alveolar proteinosis, Karagener syndrome, fibrocystic
CC pulmonary dysplasia, primary ciliary dyskinesia, pulmonary hypertension
CC and hyaline membrane disease. The present sequence is a peptide/protein
CC encoded by a single exon probe of the invention.
CC Note: The sequence data for this patent did not form part
CC of the printed specification, but was obtained in electronic
CC format directly from WIPO at
CC ftp.wipo.int/pub/published_pct_sequences.

XX SQ Sequence 56 AA;
Query Match 24.2%; Score 56; DB 23; Length 56;
Best Local Similarity 100.0%; Pred. No. 1.6e-47;
Matches 56; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 127 KIDPPVMTITQVNGSLVILHAPNLPYRYQKEKNVSIEDYVELLYRVFIINNSLEK 182
DB 1 KIDPPVMTITQVNGSLVILHAPNLPYRYQKEKNVSIEDYVELLYRVFIINNSLEK 56

RESULT 26
AAG77117
ID AAG77117 standard; Protein; 53 AA.
AC AAG77117;
XX 03-SEP-2001 (first entry)
DT Human colon cancer antigen protein SEQ ID NO:7881.
DE Human; colon cancer; colon cancer antigen; diagnosis; detection;
KW colorectal carcinoma.
XX Homo sapiens.
OS
XX WO200122920-A2.
PN
XX 05-APR-2001.
PD
XX 28-SEP-2000; 2000WO-US26524.
PF
XX 29-SEP-1999; 99US-0157137.
PR
XX 03-NOV-1999; 99US-0163280.
XX
XX (HUMA-) HUMAN GENOME SCI INC.
PA
XX Ruben SM, Barash SC, Birse CE, Rosen CA;
PI
XX WPI: 2001-235357/24.
DR
XX N-PSDB; AAH36322.
DR
XX Nucleic acids encoding 4277 human colon cancer-associated polypeptides,
PT useful for preventing, diagnosing and/or treating colorectal cancers -
PT
XX Claim 11; Page 9219-9220; 9803pp; English.
PS
XX AAH32943 to AAH37195 and AAG773514 to AAG77788 represent human colon
XX CC

CC cancer-associated nucleic acid molecules (N) and proteins (P), where
CC the proteins are collectively known as colon cancer antigens. The colon
CC cancer antigens have cytostatic activity and can be used in gene
CC therapy and vaccine production. N and P may be used in the prevention,
CC diagnosis and treatment of diseases associated with inappropriate P
CC expression. For example, N and P may be used to treat disorders
CC associated with decreased expression by rectifying mutations or deletions
CC in a patient's genome that affect the activity of P by expressing
CC inactive proteins or to supplement the patients own production of P.
CC Additionally, N may be used to produce the colon cancer-associated Ps,
CC by inserting the nucleic acids into a host cell and culturing the cell
CC to express the proteins. N and P can be used in the prevention, diagnosis
CC and treatment of colorectal carcinomas and cancers. AAH37196 to AAH37204
CC and AAH7789 represent sequences used in the exemplification of the
CC present invention.
CC N.B. Pages 666 to 682 and page 7053 of the sequence listing were
CC missing at time of publication, meaning no sequences are present for
CC SEQ ID NO:1027 to 1052, 7921 and 7922.

XX SQ Sequence 53 AA;
Query Match 3.0%; Score 7; DB 22; Length 53;
Best Local Similarity 100.0%; Pred. No. 34;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 52 ALTGNSS 58
DB 2 ALTGNSS 8

RESULT 27
AAU53889
ID AAU53889 standard; Protein; 87 AA.
XX AAU53889;
AC AAU53889;
DT 27-FEB-2002 (first entry)
DE Propionibacterium acnes immunogenic protein #14785.
XX
KW SAPHO syndrome; synovitis; acne; pustulosis; hypertosis; osteomyelitis;
KW uveitis; endophthalmitis; bone; joint; central nervous system; ELISA;
KW inflammatory lesion; acne vulgaris; enzyme linked immunosorbent assay;
KW dermatological; osteopathic; neuroprotectant.
XX
OS Propionibacterium acnes.
XX
XX WO200181581-A2.
PN
XX 01-NOV-2001.
PD
XX 20-APR-2001; 2001WO-US12865.
PF
XX 21-APR-2000; 2000US-199047P.
PR
XX 02-JUN-2000; 2000US-208841P.
PR
XX 07-JUL-2000; 2000US-216747P.
XX
XX (CORI-) CORIXA CORP.
PA
XX Skeiky YAW, Persing DH, Mitcham JL, Wang SS, Bhatia A;
PI L'maisonneuve J, Zhang Y, Jen S, Carter D;
XX
XX WPI: 2001-616774/71.
DR
XX N-PSDB; AAS59562.
DR
XX Propionibacterium acnes polypeptides and nucleic acids useful for
PT vaccinating against and diagnosing infections, especially useful for
PT treating acne vulgaris -
XX
XX Example 1; SEQ ID No 15084; 1069pp; English.
PS
XX Sequences AAU39105-AAU68017 represent Propionibacterium acnes immunogenic
XX polypeptides. The proteins and their associated DNA sequences are used in
XX CC

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OM protein - protein search, using sw model

Run on: January 13, 2003, 15:47:22 : Search time 10 Seconds

(without alignments)
448.164 Million cell updates/sec

Title: US-09-728-911-2

Perfect score: 231

Sequence: 1 MPMKHLGLISPLTGV.....YQMLDRSGRSHRCVEIP 231

Scoring table:

Gapop 60.0, Gapext 60.0

Searched: 118974 seqs, 19401057 residues

W size: 0

Total number of hits satisfying chosen parameters: 118974

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 100 summaries

Database:

Published Applications AA:
1: /cgn2_6/ptodata/1/pubppaa/US08_NEW_PUB.pep:*
2: /cgn2_6/ptodata/1/pubppaa/PCR_NEW_PUB.pep:*
3: /cgn2_6/ptodata/1/pubppaa/US06_NEW_PUB.pep:*
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5: /cgn2_6/ptodata/1/pubppaa/US07_NEW_PUB.pep:*
6: /cgn2_6/ptodata/1/pubppaa/US07_PUBCOMB.pep:*
7: /cgn2_6/ptodata/1/pubppaa/US08_PUBCOMB.pep:*
8: /cgn2_6/ptodata/1/pubppaa/US08_PUBCOMB.pep:*
9: /cgn2_6/ptodata/1/pubppaa/US09_NEW_PUB.pep:*
10: /cgn2_6/ptodata/1/pubppaa/US09_PUBCOMB.pep:*
11: /cgn2_6/ptodata/1/pubppaa/US10_NEW_PUB.pep:*
12: /cgn2_6/ptodata/1/pubppaa/US10_PUBCOMB.pep:*
13: /cgn2_6/ptodata/1/pubppaa/US60_NEW_PUB.pep:*
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	231	100.0	231	10	US-09-728-911-2
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3	210	90.9	210	10	US-09-728-911-13
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93 Sequence 2, Appli
94 Sequence 6104, Ap
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96 Sequence 11, Appl
97 Sequence 12, Appl
98 Sequence 17, Appl
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6 2.6 458 9 US-09-738-626-6104
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ALIGNMENTS

RESULT 1
US-09-728-911-2
; Sequence 2, Application US/09728911
; Patent No. US20020012669A1
; GENERAL INFORMATION:
; APPLICANT: Presnell, Scott R.
; APPLICANT: Xu, Wenfeng
; APPLICANT: Kindsvogel, Wayne
; APPLICANT: Chen, Zhi
; TITLE OF INVENTION: Human Cytokine Receptor
; FILE REFERENCE: 99-93
; CURRENT APPLICATION NUMBER: US/09/728,911
; PRIOR FILING DATE: 2000-12-01
; PRIOR APPLICATION NUMBER: US 60/169,049
; PRIOR FILING DATE: 1999-12-03
; PRIOR APPLICATION NUMBER: US 60/232,219
; PRIOR FILING DATE: 2000-09-13
; PRIOR APPLICATION NUMBER: US 60/244,610
; PRIOR FILING DATE: 2000-10-31
; NUMBER OF SEQ ID NOS: 36
; SOFTWARE: FastSeq for Windows Version 3.0
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; ORGANISM: Homo sapiens
US-09-728-911-2

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; Sequence 6, Application US/09949192
; Patent No. US20020142292A1
; GENERAL INFORMATION:
; APPLICANT: Patham, Christi L.
; APPLICANT: Gorman, Daniel L.
; APPLICANT: Kurata, Hirokazu
; APPLICANT: Arai, Naoko
; APPLICANT: Sana, Theodore R.
; APPLICANT: Mattson, Jeanine D.
; APPLICANT: Murphy, Erin E.

; APPLICANT: Savkoor, Chetan
; APPLICANT: Grein, Jeffery
; APPLICANT: Smith, Kathleen M.
; APPLICANT: McClanahan, Terrill K.
; TITLE OF INVENTION: MAMMALIAN GENES; RELATED REAGENTS AND METHODS
; FILE REFERENCE: DX01169K
; CURRENT APPLICATION NUMBER: US/09/949,192
; CURRENT FILING DATE: 2001-09-07
; PRIOR APPLICATION NUMBER: 60/231,267
; PRIOR FILING DATE: 2000-09-08
; NUMBER OF SEQ ID NOS: 53
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 6
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US-09-949-192-6

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; Patent No. US20020012669A1
; GENERAL INFORMATION:
; APPLICANT: Presnell, Scott R.
; APPLICANT: Xu, Wenfeng
; APPLICANT: Kindsvogel, Wayne
; APPLICANT: Chen, Zhi
; TITLE OF INVENTION: Human Cytokine Receptor
; FILE REFERENCE: 99-93
; CURRENT APPLICATION NUMBER: US/09/728,911
; CURRENT FILING DATE: 2000-12-01
; PRIOR APPLICATION NUMBER: US 60/169,049
; PRIOR FILING DATE: 1999-12-03
; PRIOR APPLICATION NUMBER: US 60/232,219
; PRIOR FILING DATE: 2000-09-13
; PRIOR APPLICATION NUMBER: US 60/244,610
; PRIOR FILING DATE: 2000-10-31
; NUMBER OF SEQ ID NOS: 36
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; SEQ ID NO 13
; LENGTH: 210
; TYPE: PRT
; ORGANISM: Homo sapiens
US-09-728-911-13

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Pred. No. is the number of results predicted by chance to have a
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and is derived by analysis of the total score distribution.

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ALIGNMENTS

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; Patent No. 5177189
; APPLICANT: DYER, CHERYL A.; CURTISS, LINDA K.; SMITH, RICHARD
; TITLE OF INVENTION: POLYPEPTIDE ANALOGS OF APOLIPOPROTEIN E
; NUMBER OF SEQUENCES: 11
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/07/395,732
; FILING DATE: 18-AUG-1989
; SEQ ID NO: 11:
; LENGTH: 30
5177189-11

Query Match 2.6%; Score 6; DB 6; Length 30;
Best Local Similarity 100.0%; Pred. No. 32;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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DB 15 GRVRAA 20

RESULT 2

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; Patent No. 5182364
; APPLICANT: DYER, CHERYL A.; CURTISS, LINDA K.; SMITH, RICHARD
; TITLE OF INVENTION: POLYPEPTIDE ANALOGS OF APOLIPOPROTEIN E
; NUMBER OF SEQUENCES: 14
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/485,158
; FILING DATE: 26-FEB-1990
; SEQ ID NO: 12:
; LENGTH: 30
5182364-12

Query Match 2.6%; Score 6; DB 6; Length 30;
Best Local Similarity 100.0%; Pred. No. 32;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 101 GRVRAA 106
DB 15 GRVRAA 20

RESULT 3

US-08-023-980B-36
; Sequence 36, Application US/08023980B
; Patent No. 5843641
; GENERAL INFORMATION:
; APPLICANT: Brown, Robert
; APPLICANT: Horvitz, H. Robert
; APPLICANT: Rosen, Daniel R.
; TITLE OF INVENTION: COMPOUNDS AND METHODS FOR THE DIAGNOSIS,
; TREATMENT AND PREVENTION OF DISEASES OF CELL DEATH
; NUMBER OF SEQUENCES: 45
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Clark & Elbing LLP
; STREET: 585 Commercial Street
; CITY: Boston
; STATE: MA
; COUNTRY: USA
; ZIP: 02109-1024
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patent In Release #1.0, Version #1.30
; CURRENT APPLICATION DATA:

; APPLICATION NUMBER: US/08/023,980B
; FILING DATE: 26-FEB-1993
; CLASSIFICATION: 435
; ATTORNEY/AGENT INFORMATION:
; NAME: Clark, Paul T.
; REGISTRATION NUMBER: 30,162
; REFERENCE/DOCKET NUMBER: 00786/177001
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617/723-4123
; TELEFAX: 617/723-8962
; TELEX:
; INFORMATION FOR SEQ ID NO: 36:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 39 amino acids
; TYPE: amino acid
; STRANDEDNESS: not relevant
; TOPOLOGY: linear
; MOLECULE TYPE: protein
US-08-023-980B-36

Query Match 2.6%; Score 6; DB 2; Length 39;
Best Local Similarity 100.0%; Pred. No. 41;
Matches 6; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 160 NVSIED 165
DB 17 NVSIED 22

RESULT 4

US-08-486-953A-31
; Sequence 31, Application US/08486953A
; Patent No. 5849290
; GENERAL INFORMATION:
; APPLICANT: Brown, Robert
; APPLICANT: Horvitz, H. Robert
; APPLICANT: Rosen, Daniel R.
; TITLE OF INVENTION: COMPOUNDS AND METHODS FOR THE DIAGNOSIS,
; TREATMENT AND PREVENTION OF DISEASES OF CELL DEATH
; NUMBER OF SEQUENCES: 53
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Clark & Elbing LLP
; STREET: 176 Federal Street
; CITY: Boston
; STATE: MA
; COUNTRY: USA
; ZIP: 02110
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: FastSeq
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/486,953A
; FILING DATE: 07-JUN-1995
; CLASSIFICATION: 424
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: US 08/204,052
; FILING DATE: 28-FEB-1994
; ATTORNEY/AGENT INFORMATION:
; NAME: Clark, Paul T.
; REGISTRATION NUMBER: 30,162
; REFERENCE/DOCKET NUMBER: 00786/223002
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: 617/428-0200
; TELEFAX: 617/428-7045
; TELEX:
; INFORMATION FOR SEQ ID NO: 31:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 39 amino acids
; TYPE: amino acid
; STRANDEDNESS: not relevant
; TOPOLOGY: linear

GenCore version 5.1.3
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OM protein - protein search, using sw model

Run on: January 13, 2003, 15:44:16 ; Search time 34 Seconds
(without alignments)
1399,908 Million cell updates/sec

Title: US-09-728-911-2

Perfect score: 231
Sequence: 1 MPMKCTFLGFLISFLLTGA.....YQMLDRRSQSRSERCVEIP 231

Scoring table: OLIGO

Gapop 60.0, Gapext 60.0

Searched: 671580 seqs, 206047115 residues

W size: 0

Total number of hits satisfying chosen parameters: 671580

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 100 summaries

Database:

1: SP archaea: *
2: SP bacteria: *
3: SP fungi: *
4: SP human: *
5: SP invertebrate: *
6: SP mammal: *
7: SP mhc: *
8: SP organelle: *
9: SP phage: *
10: SP plant: *
11: SP rodent: *
12: SP virus: *
13: SP vertebrate: *
14: SP unclassified: *
15: SP viirus: *
16: SP bacteriophage: *
17: SP archaea: *

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	231	100.0	231	4 Q96A41	Q96A41 homo sapien
2	165	71.4	263	4 Q969J5	Q969J5 homo sapien
3	125	54.1	130	4 Q96OR0	Q96OR0 homo sapien
4	8	3.5	121	17 Q9HKC4	Q9HKC4 thermoplasma
5	8	3.5	698	11 Q9R3J2	Q9R3J2 mus musculu
6	8	3.5	699	11 Q9D4A6	Q9D4A6 mus musculu
7	8	3.5	833	16 Q8ZKM7	Q8ZKM7 salmoneila
8	7	3.0	83	17 Q8TUL7	Q8TUL7 methanosarc
9	7	3.0	127	17 Q8TMM9	Q8TMM9 methanosarc
10	7	3.0	148	16 Q8XPS5	Q8XPS5 clostridium
11	7	3.0	180	13 Q8UW74	Q8UW74 xenopus lae
12	7	3.0	180	13 Q8UW73	Q8UW73 xenopus lae
13	7	3.0	180	13 Q8UW72	Q8UW72 xenopus lae
14	7	3.0	181	5 Q9GV59	Q9GV59 drosophila
15	7	3.0	192	8 Q63049	Q63049 rhizogonium
16	7	3.0	199	2 Q9BRX7	Q9BRX7 corynebacte

17	7	3.0	213	2 Q93MY6	Q93MY6 lactococcus
18	7	3.0	218	16 Q9CES2	Q9CES2 lactococcus
19	7	3.0	218	10 Q9LY15	Q9LY15 arabidopsis
20	7	3.0	225	11 Q9Z1B2	Q9Z1B2 rattus norv
21	7	3.0	231	8 Q8WA03	Q8WA03 laticauda c
22	7	3.0	236	16 Q8UBG5	Q8UBG5 agrobacteri
23	7	3.0	241	16 Q8XU48	Q8XU48 rhizobium m
24	7	3.0	243	16 Q8UG20	Q8UG20 agrobacteri
25	7	3.0	255	16 Q9X015	Q9X015 thermotoga
26	7	3.0	303	16 Q8XBX7	Q8XBX7 escherichia
27	7	3.0	304	5 Q46594	Q46594 caenorhabdi
28	7	3.0	312	16 Q9X7V5	Q9X7V5 streptomyce
29	7	3.0	333	17 Q980J8	Q980J8 sulfolobus
30	7	3.0	348	5 Q95W37	Q95W37 plasmidium
31	7	3.0	357	16 Q66804	Q66804 aquifex aeo
32	7	3.0	359	16 Q9CK69	Q9CK69 pasteurella
33	7	3.0	367	17 Q8TT19	Q8TT19 methanosarc
34	7	3.0	386	8 Q8SM15	Q8SM15 stigeocloni
35	7	3.0	393	16 Q8URK1	Q8URK1 agrobacteri
36	7	3.0	393	16 Q8UDU0	Q8UDU0 agrobacteri
37	7	3.0	407	16 Q931X6	Q931X6 streptomyce
38	7	3.0	434	17 Q9HJ87	Q9HJ87 thermoplasma
39	7	3.0	439	17 Q97BG4	Q97BG4 thermoplasma
40	7	3.0	446	10 Q98RQ9	Q98RQ9 guillardia
41	7	3.0	448	16 Q97L22	Q97L22 clostridium
42	7	3.0	479	10 Q9L207	Q9L207 arabidopsis
43	7	3.0	489	16 Q985L6	Q985L6 rhizobium 1
44	7	3.0	495	5 Q9VHX1	Q9VHX1 drosophila
45	7	3.0	512	17 Q26458	Q26458 methanobact
46	7	3.0	552	10 Q957U1	Q957U1 arabidopsis
47	7	3.0	553	3 Q94155	Q94155 pichia stip
48	7	3.0	562	17 Q9YCC5	Q9YCC5 aeropyrum p
49	7	3.0	595	10 Q9CSF4	Q9CSF4 arabidopsis
50	7	3.0	637	16 Q92J39	Q92J39 rickettsia
51	7	3.0	788	5 Q9XWE1	Q9XWE1 caenorhabdi
52	7	3.0	800	13 Q91551	Q91551 xenopus lae
53	7	3.0	823	16 Q8YHC6	Q8YHC6 brucella me
54	7	3.0	921	16 Q9ZD78	Q9ZD78 rickettsia
55	7	3.0	929	6 Q9BDT5	Q9BDT5 rhychoecyon
56	7	3.0	960	16 Q8Y377	Q8Y377 ralistonia s
57	7	3.0	1041	2 Q93C90	Q93C90 actinomadr
58	7	3.0	1062	2 Q9RC22	Q9RC22 bacillus sp
59	7	3.0	1064	16 Q8XEP1	Q8XEP1 ralistonia s
60	7	3.0	1210	13 Q92137	Q92137 xenopus lae
61	7	3.0	1291	2 Q93H21	Q93H21 streptomyce
62	7	3.0	1385	5 Q8WT26	Q8WT26 leishmania
63	7	3.0	2771	11 Q9WTS7	Q9WTS7 mus musculu
64	7	3.0	2825	11 Q70465	Q70465 mus musculu
65	7	2.6	21	6 Q9TRC5	Q9TRC5 canis fami1
66	7	2.6	31	2 Q45547	Q45547 bacillus su
67	6	2.6	31	2 Q54825	Q54825 streptococc
68	6	2.6	54	9 Q9A2P4	Q9A2P4 bacterioph
69	6	2.6	56	16 Q97RW6	Q97RW6 streptococc
70	6	2.6	57	16 Q8UC09	Q8UC09 agrobacteri
71	6	2.6	63	2 Q85792	Q85792 enterococcu
72	6	2.6	68	17 Q981M8	Q981M8 rhizobium 1
73	6	2.6	71	17 Q8TKS8	Q8TKS8 methanosarc
74	6	2.6	73	10 Q8W2M4	Q8W2M4 oryza sativ
75	6	2.6	79	2 P72499	P72499 streptococc
76	6	2.6	80	2 Q70028	Q70028 streptomyce
77	6	2.6	81	2 Q93SG3	Q93SG3 eubacterium
78	6	2.6	82	16 Q50403	Q50403 mycobacteri
79	6	2.6	84	5 Q9VCK3	Q9VCK3 drosophila
80	6	2.6	87	13 Q91876	Q91876 xenopus ami
81	6	2.6	87	13 Q91875	Q91875 xenopus ruw
82	6	2.6	87	13 Q91BB4	Q91BB4 xenopus mue
83	6	2.6	87	13 Q91BB7	Q91BB7 xenopus lae
84	6	2.6	87	13 Q91BC3	Q91BC3 xenopus lae
85	6	2.6	87	13 Q91BB8	Q91BB8 xenopus lae
86	6	2.6	87	13 Q91BC2	Q91BC2 xenopus lae
87	6	2.6	87	13 Q91BC1	Q91BC1 xenopus lae
88	6	2.6	87	13 Q91BB6	Q91BB6 xenopus bor
89	6	2.6	87	13 Q91BB6	Q91BB6 xenopus bor

90 Q91bb3 xenopus ves
91 Q91bb1 xenopus ruw
92 Q91bb0 xenopus tro
93 Q93kq8 yersinia en
94 Q97qk7 streptococc
95 Q9vsg9 drosophila
96 Q8x1y0 mus musculu
97 Q9pp75 campylobact
98 Q9f7z6 erwinia amy
99 Q41409 human immun
100 Q82z58 salmonella

ALIGNMENTS

RESULT 1
Q96A41 ID Q96A41 PRELIMINARY; PRT; 231 AA.
AC Q96A41, 231 (TREMBLrel. 19, Created)
DT 01-DEC-2001 (TREMBLrel. 19, Last sequence update)
DT 01-MAR-2002 (TREMBLrel. 20, Last annotation update)
DE Soluble cytokine class II receptor, short isoform precursor
DE (Interleukin 22-binding protein CRF2-10) (Class II cytokine receptor)
DE (Interleukin-22 binding protein)
GN CRF2-S1 OR IL22BP OR IL22RA2 OR IL-22BP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=MAMMARY GLAND;
RX MEDLINE=21518574; PubMed=11607789;
RA Gruenberg B.H., Schoenemeyer A., Weiss B., Toschi L., Kunz S.,
RA Wolk K., Asadullah K., Sabat R.;
RT "A novel, soluble homologue of the human IL-10 receptor with
RT preferential expression in placenta.";
RL Genes Immun. 2:329-334(2001).
RN [2]
RP SEQUENCE FROM N.A.
RX MEDLINE=21286453; PubMed=11390454;
RA Kottenko S.V., Izotova L.S., Mirochnitchenko O.V., Esterova E.,
RA Chen Z., Dillon S.R., Gao Z., Gilbert T., Madden K., Schlutsmeyer S.,
RA Yao L., Whitmore T.E., Chandrasekhar Y., Grant F.J., Maurer M.,
RA Jelinek L., Storey H., Brender T., Hammond A., Topouzis S.,
RA Clegg C.H., Foster D.C.;
RT "A soluble class II cytokine receptor, IL-22RA2, is a naturally
RT occurring IL-22 antagonist.";
RL Proc. Natl. Acad. Sci. U.S.A. 98:9511-9516(2001).
RN [4]
RP SEQUENCE FROM N.A.
RC TISSUE=BREAST;
RA Dumoutier L., Lejeune D., Renaud J.C.;
RT "Cloning and characterization of Interleukin-22 Binding Protein (IL-
RT 22BP), a natural antagonist of IL-TIF/IL-22.";
RL Submitted (DEC-2000) to the EMBL/GenBank/DBJ databases.
DR EMBL; AJ313161; CAC85634.1; -
DR EMBL; AY040566; AAK85714.1; -
DR EMBL; AY044429; AAK91775.1; -
DR EMBL; AJ297262; CAC83097.1; -
DR InterPro; IPR000282; Cytok_receptor_2.
KW Receptor; Signal.
FT SIGNAL 1 21 POTENTIAL.

FT CHAIN 22 231 SOLUBLE CYTOKINE CLASS II RECEPTOR, SHORT
FT ISOFORM.
SQ SEQUENCE 231 AA; 26979 MW; 24A6912BFF75100F CRC64;
Query Match 100.0%; Score 231; DB 4; Length 231;
Best Local Similarity 100.0%; Pred. No. 3.1e-234;
Matches 231; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 MNPKECFGLISFLTGVAGTQSTHESLKQPVQFQSRNFHNLQWOPGRLTGNSSVY 60
DB 1 MNPKECFGLISFLTGVAGTQSTHESLKQPVQFQSRNFHNLQWOPGRLTGNSSVY 60
QY 61 FVQYKIYGORQWKNEKDCWGTQELSCDLTSETSDIQEYVYGRVRAASAGSYSEWSMTFRF 120
DB 61 FVQYKIYGORQWKNEKDCWGTQELSCDLTSETSDIQEYVYGRVRAASAGSYSEWSMTFRF 120
QY 121 TPWETKIDPPVMNITQVNGSLLVILHAPNLPYRYQKEKNVSIEDYVYLLYRVFIINNSL 180
DB 121 TPWETKIDPPVMNITQVNGSLLVILHAPNLPYRYQKEKNVSIEDYVYLLYRVFIINNSL 180
QY 181 EXEQVYEGAHRAVEIEALTPHSSYCVVAEYIYQPMIDRRSORSERCVIEIP 231
DB 181 EXEQVYEGAHRAVEIEALTPHSSYCVVAEYIYQPMIDRRSORSERCVIEIP 231
RESULT 2
Q969J5 ID Q969J5 PRELIMINARY; PRT; 263 AA.
AC Q969J5;
DT 01-DEC-2001 (TREMBLrel. 19, Created)
DT 01-DEC-2001 (TREMBLrel. 19, Last sequence update)
DT 01-MAR-2002 (TREMBLrel. 20, Last annotation update)
DE Soluble cytokine class II receptor, long isoform precursor
DE (Interleukin 22-binding protein CRF2-10L).
GN CRF2-S1 OR IL22BP.
OS Homo sapiens (Human).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX NCBI_TaxID=9606;
RN [1]
RP SEQUENCE FROM N.A.
RC TISSUE=PLACENTA;
RX MEDLINE=21518574; PubMed=11607789;
RA Gruenberg B.H., Schoenemeyer A., Weiss B., Toschi L., Kunz S.,
RA Wolk K., Asadullah K., Sabat R.;
RT "A novel, soluble homologue of the human IL-10 receptor with
RT preferential expression in placenta.";
RL Genes Immun. 2:329-334(2001).
RN [2]
RP SEQUENCE FROM N.A.
RX MEDLINE=21286453; PubMed=11390454;
RA Kottenko S.V., Izotova L.S., Mirochnitchenko O.V., Esterova E.,
RA Dickensheets H., Donnelly R.P., Pestka S.;
RT "Identification, cloning, and characterization of a novel soluble
RT receptor that binds IL-22 and neutralizes its activity.";
RL J. Immunol. 166:7096-7103(2001).
DR EMBL; AJ313162; CAC85635.1; -
DR EMBL; AY040567; AAK85715.1; -
DR InterPro; IPR000282; Cytok_receptor_2.
KW Receptor; Signal.
FT SIGNAL 1 21 POTENTIAL.
FT CHAIN 22 263 SOLUBLE CYTOKINE CLASS II RECEPTOR, LONG
FT ISOFORM.
SQ SEQUENCE 263 AA; 30550 MW; C96EBC5D78AC79B CRC64;
Query Match 71.4%; Score 165; DB 4; Length 263;
Best Local Similarity 100.0%; Pred. No. 7.5e-165;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 67 YGQRQWKNEKDCWGTQELSCDLTSETSDIQEYVYGRVRAASAGSYSEWSMTFRFPWNET 126
DB 99 YGQRQWKNEKDCWGTQELSCDLTSETSDIQEYVYGRVRAASAGSYSEWSMTFRFPWNET 158

QY 127 KIDPVMNITOVNGSLVTLHAHNLPRYOKKKNVSIIDYELLYRVFIINNSLEKEQKV 186
 DB 159 KIDPVMNITOVNGSLVTLHAHNLPRYOKKKNVSIIDYELLYRVFIINNSLEKEQKV 218
 QY 187 YEGAHRAVEIEALTPHSSCYVAETIQPMLDRSRORSERCEIP 221
 DB 219 YEGAHRAVEIEALTPHSSCYVAETIQPMLDRSRORSERCEIP 263

RESULT 3

Q96OR0 PRELIMINARY; PRT; 130 AA.
 AC Q96OR0;
 DT 01-DEC-2001 (TREMBlrel. 19, Created)
 DT 01-DEC-2001 (TREMBlrel. 19, Last sequence update)
 DT 01-MAR-2002 (TREMBlrel. 20, Last annotation update)
 DE Interleukin 22-binding protein CRP2-10S.
 GN IL22BP.
 OS Homo sapiens (Human).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.
 NCBI_TaxID=9606;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX MEDLINE=21286453; PubMed=11390454;
 RA Korenko S.V., Izotova L.S., Mitrochitschenko O.V., Esterova E.,
 RA Dickensheats H., Donnelly R.P., Pestka S.,
 RT "Identification, cloning, and characterization of a novel soluble
 receptor that binds IL-22 and neutralizes its activity.",
 RL J. Immunol. 166:7096-7103(2001).
 DR EMBL; AY040568; AAK85716.1; -
 DR InterPro; IPR000282; Cytok_receptor 2.
 SQ SEQUENCE 130 AA; 15128 MW; A165814C641F5E5B CRC64;

Query Match 54.1%; Score 125; DB 4; Length 130;
 Best Local Similarity 100.0%; Pred. No. 4.5e-123;
 Matches 125; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 MWPKRCFLGFLISFLTVAGTOSTHESLKPQVQFQSRNFHNTLQWPGRALTGNSSVY 60
 DB 1 MWPKRCFLGFLISFLTVAGTOSTHESLKPQVQFQSRNFHNTLQWPGRALTGNSSVY 60
 QY 61 FVQYKIVGQKQKEDCMGTQELSCDLTSETSDIOEPYGRVAAAGSYSEMSWTPPR 120
 DB 61 FVQYKIVGQKQKEDCMGTQELSCDLTSETSDIOEPYGRVAAAGSYSEMSWTPPR 120
 QY 121 TPWME 125
 DB 121 TPWME 125

RESULT 4

Q9HKC4 PRELIMINARY; PRT; 121 AA.
 AC Q9HKC4;
 DT 01-MAR-2001 (TREMBlrel. 16, Created)
 DT 01-MAR-2001 (TREMBlrel. 16, Last sequence update)
 DT 01-MAR-2001 (TREMBlrel. 16, Last annotation update)
 DE Hypothetical membrane protein.
 GN TA0677.
 OS Thermoplasma acidophilum.
 OC Archaea; Euryarchaeota; Thermoplasmatia; Thermoplasmales;
 OC Thermoplasmataceae; Thermoplasma.
 NCBI_TaxID=2303;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX STRAIN=DSM 1728;
 RX MEDLINE=20479972; PubMed=11029001;
 RA Rupp A., Graml W., Santos-Martinez M.-L., Koretke K.K., Volker C.,
 RA Mewes H.-W., Frishman D., Stocker S., Lups A.N., Baumeister W.,
 RT "The genome sequence of the thermophilic scavenger Thermoplasma
 Nature 407:508-513(2000)."

DR EMBL; AL445065; CAC11815.1; -
 KW Hypothetical protein; Complete proteome.
 SQ SEQUENCE 121 AA; 13083 MW; 3C80DC4C2B04FD39 CRC64;

Query Match

Best Local Similarity 3.5%; Score 8; DB 17; Length 121;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 139 NSGLVLVL 146
 DB 77 NSGLVLVL 84

RESULT 5

Q8R322 PRELIMINARY; PRT; 698 AA.
 AC Q8R322;
 DT 01-JUN-2002 (TREMBlrel. 21, Created)
 DT 01-JUN-2002 (TREMBlrel. 21, Last sequence update)
 DT 01-JUN-2002 (TREMBlrel. 21, Last annotation update)
 DE Similar to RIKEN cDNA 4933405K21 gene.
 OS Mus musculus (Mouse).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 NCBI_TaxID=10090;
 RN [1]
 RP SEQUENCE FROM N.A.
 RA Strausberg R.,
 RL Submitted (APR-2002) to the EMBL/GenBank/DBJ databases.
 DR EMBL; BC026797; AAH26797.1; -
 SQ SEQUENCE 698 AA; 79560 MW; 22E5E7217AA4D33 CRC64;

Query Match 3.5%; Score 8; DB 11; Length 698;
 Best Local Similarity 100.0%; Pred. No. 15;
 Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 21 GTQSTHES 28
 DB 106 GTQSTHES 113

RESULT 6

Q9D4A6 PRELIMINARY; PRT; 699 AA.
 AC Q9D4A6;
 DT 01-JUN-2001 (TREMBlrel. 17, Created)
 DT 01-JUN-2001 (TREMBlrel. 17, Last sequence update)
 DT 01-JUN-2001 (TREMBlrel. 17, Last annotation update)
 DE 4933405K21RIK protein.
 GN 4933405K21RIK.
 OS Mus musculus (Mouse).
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
 NCBI_TaxID=10090;
 RN [1]
 RP SEQUENCE FROM N.A.
 RX STRAIN=C57BL/6J; TISSUE=TESTIS;
 RX MEDLINE=21085660; PubMed=11217851;
 RA Kawai J., Shingawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
 RA Akawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
 RA Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
 RA Saito T., Okazaki Y., Gojobori T., Bono H., Kasukawa T., Saito R.,
 RA Kadota K., Matsuda H.A., Ashburner M., Batilov S., Casavant T.,
 RA Fleischmann W., Gaasterland T., Gissi C., King B., Kochwa H.,
 RA Kuehl P., Lewis S., Matsuo Y., Nikaido I., Pesole G., Quackenbush J.,
 RA Schirimi L.M., Staudt F., Suzuki R., Tomita M., Wagner L., Washio T.,
 RA Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
 RA Blake J., Boilelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
 RA Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
 RA Gustinich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
 RA Lyons P., Marchionni L., Mashima J., Mazzarelli U., Mommaerts P.,
 RA Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
 RA Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,

RA Suzuki H., Toyooka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
RA Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA Hayashizaki Y.,
RA "Functional annotation of a full-length mouse cDNA collection.";
RL Nature 409:685-690(2001).
DR EMBL; AK016671; BAB30371.1; -.
DR MGD; MGI:1921662; 4933405K21Rik.
SQ SEQUENCE 699 AA; 79574 MW; 5403C38F9DC7ED75 CRC64;

Query Match 3.5%; Score 8; DB 11; Length 699;
Best Local Similarity 100.0%; Pred. No. 15;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 21 GTQSTHES 28
|||
DB 107 GTQSTHES 114

RESULT 7
Q8ZKM7 PRELIMINARY; PRT; 833 AA.
AC Q8ZKM7
DT 01-MAR-2002 (TrEMBLrel. 20, Created)
DT 01-MAR-2002 (TrEMBLrel. 20, Last sequence update)
DT 01-JUN-2002 (TrEMBLrel. 21, Last annotation update)
DE General PTS family, enzyme I (EC 2.7.3.9).
DE PTSA OR STM4110.
GN Salmonella typhimurium.
OS Bacteria; Proteobacteria; gamma subdivision; Enterobacteriaceae;
OC Bacteria.
OC Salmonella.
NCBI_TaxID=602;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=LT2 / SGSC1412 / ATCC 700720;
RX MEDLINE=21534948; PubMed=11677609;
RA McClelland M., Sanderson K.E., Spieth J., Clifton S.W., Latreille P.,
RA Courtney L., Porwollik S., Ali J., Dante M., Du F., Hou S., Layman D.,
RA Leonard S., Nguyen C., Scott K., Holmes A., Grewal N., Mulvaney E.,
RA Ryan E., Sun H., Florea L., Miller W., Stoneking T., Nhan M.,
RA Waterston R., Wilson R.K.;
RA "Complete genome sequence of Salmonella enterica serovar Typhimurium
LT2.";
RL Nature 413:852-856(2001).
DR EMBL; AE008892; AAL22950.1; -.
DR InterPro; IPR001020; HPr_Hisp_site.
DR InterPro; IPR000032; HPr_protein.
DR InterPro; IPR000121; PEP_utilizers.
DR InterPro; IPR004715; PTSIIA fruc.
DR InterPro; IPR002178; PTS_EIIA_2.
DR Pfam; PF00391; PEP-utilizers; 1.
DR Pfam; PF02896; PEP-utilizers_C; 1.
DR Pfam; PF00381; PRS-HPr; 1.
DR Pfam; PF00359; PTS_EIIA_2; 1.
DR ProDom; PD000940; PEP_utilizers; 1.
DR ProDom; PD001689; PTS_EIIA_2; 1.
DR TIGRFAMs; TIGR00848; frua; 1.
DR PROSITE; PS00742; PEP_ENZYMES_2; 1.
DR PROSITE; PS00370; PEP_ENZYMES_PHOS_SITE; 1.
DR PROSITE; PS00369; PTS_HPR_His; 1.
KW Transference; Complete proteome.
SQ SEQUENCE 833 AA; 92082 MW; D2FE53D3DD81D6BF CRC64;

Query Match 3.5%; Score 8; DB 16; Length 833;
Best Local Similarity 100.0%; Pred. No. 17;
Matches 8; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 194 VEIEALTP 201
|||||
DB 317 VEIEALTP 324

RESULT 8
O8TUL7

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OM protein - protein search, using sw model

Run on: January 13, 2003, 15:43:26 ; Search time 11 Seconds

(without alignments)
871,003 Million cell updates/sec

Title: US-09-728-911-2
Perfect score: 231
Sequence: 1 MPMKHCFLGFLISFLLTVA.....YQPMIDRRSQRSEKCEVLP 231

Scoring table: OLIGO
Gapop 60.0 , Gapext 60.0

Searched: 112892 seqs, 41476328 residues

Wc size: 0

Total number of hits satisfying chosen parameters: 112892

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Listing first 100 summaries

Database: SwissProt_40:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	7	3.0	208	1	HIS1_LACLA
2	7	3.0	213	1	HIS1_LEGPN
3	7	3.0	270	1	CX83_HUMAN
4	7	3.0	297	1	RLA0_METVO
5	7	3.0	303	1	TTDA_ECOLI
6	7	3.0	341	1	RIR2_HELPY
7	7	3.0	457	1	YIRO_YEAST
8	7	3.0	535	1	ORC2_SCHPO
9	7	3.0	546	1	GHT5_SCHPO
10	7	3.0	567	1	HXT9_YEAST
11	7	3.0	567	1	HXT9_YEAST
12	7	3.0	570	1	HXT1_YEAST
13	7	3.0	592	1	HXT5_YEAST
14	7	3.0	637	1	FTSH_RICCN
15	7	3.0	637	1	FTSH_RICCN
16	7	3.0	812	1	FTSH_RICCN
17	7	3.0	1166	1	LPK2_EUPOC
18	7	3.0	4273	1	PKGM_BACSU
19	6	2.6	65	1	BR12_SCHCO
20	6	2.6	91	1	Y022_ARCFU
21	6	2.6	119	1	EYAI_CHICK
22	6	2.6	119	1	EYAI_CHICK
23	6	2.6	125	1	YSH6_CAEEL
24	6	2.6	128	1	YF75_MYCPN
25	6	2.6	134	1	CYB_ANOCU
26	6	2.6	134	1	CYB_STUTI
27	6	2.6	139	1	CDBB_CLOAB
28	6	2.6	144	1	NU6M_ASCSU
29	6	2.6	145	1	PSAN_HORVU
30	6	2.6	152	1	SODC_CAVPO
31	6	2.6	153	1	SODC_MOUSE
32	6	2.6	153	1	SODC_RAT
33	6	2.6	157	1	NU5B_XYLFRA

34	6	2.6	161	1	PHAB_SYNEL
35	6	2.6	161	1	PHAB_SYNY3
36	6	2.6	161	1	PHAB_SYNY4
37	6	2.6	162	1	PHAB_PREDI
38	6	2.6	166	1	MEG3_ARATH
39	6	2.6	176	1	CYB_NYCHU
40	6	2.6	176	1	CYB_STULI
41	6	2.6	182	1	PGRE_TRINI
42	6	2.6	185	1	BCNA_CLOPE
43	6	2.6	195	1	OA23_MOUSE
44	6	2.6	203	1	RECR_MYCLE
45	6	2.6	204	1	YAGU_ECOLI
46	6	2.6	207	1	GSHP_AERHY
47	6	2.6	215	1	KAD_MYCPU
48	6	2.6	216	1	HIS2_THIEET
49	6	2.6	219	1	GSHE_PIG
50	6	2.6	220	1	RECR_DEIRA
51	6	2.6	221	1	GSHE_CANFA
52	6	2.6	221	1	GSHE_HUMAN
53	6	2.6	221	1	GSHE_HUMAN
54	6	2.6	230	1	RNS1_ARATH
55	6	2.6	232	1	SOML_PROAN
56	6	2.6	234	1	DNAJ_RHILE
57	6	2.6	234	1	PUR7_PYPAR
58	6	2.6	234	1	YRUE_LACLA
59	6	2.6	239	1	FRDB_WOLSU
60	6	2.6	243	1	ZIPA_XYLFRA
61	6	2.6	244	1	PHOS_MOUSE
62	6	2.6	246	1	PHOS_RAT
63	6	2.6	254	1	YFA6_YEAST
64	6	2.6	255	1	TPIS_SALTI
65	6	2.6	255	1	TPIS_SALTI
66	6	2.6	259	1	CFAD_MOUSE
67	6	2.6	259	1	DERP3_DERPA
68	6	2.6	259	1	Y305_CHLMU
69	6	2.6	271	1	PSB8_RAT
70	6	2.6	272	1	ESL3_MYCPN
71	6	2.6	278	1	PSB8_HUMAN
72	6	2.6	280	1	P29K_STRPN
73	6	2.6	280	1	P29K_STRPN
74	6	2.6	285	1	CABA_MOUSE
75	6	2.6	285	1	YOV9_CAEEL
76	6	2.6	289	1	CWPN_SCHPO
77	6	2.6	289	1	TF_CAVPO
78	6	2.6	290	1	NIH2_AZOCH
79	6	2.6	292	1	FIXA_RHINE
80	6	2.6	294	1	NIFH_BRATA
81	6	2.6	294	1	NIFH_BRATA
82	6	2.6	297	1	YG1B_YEAST
83	6	2.6	299	1	PUR7_STRCO
84	6	2.6	302	1	POB4_FOMPV
85	6	2.6	302	1	POB4_FOMPV
86	6	2.6	305	1	POB4_PSEET
87	6	2.6	305	1	CDSA_MYGCE
88	6	2.6	305	1	TNE2_HUMAN
89	6	2.6	309	1	KHSE_BUCAI
90	6	2.6	317	1	APB_HUMAN
91	6	2.6	317	1	APB_HUMAN
92	6	2.6	317	1	APB_HUMAN
93	6	2.6	322	1	UNG2_MOUSE
94	6	2.6	326	1	UNG2_MOUSE
95	6	2.6	329	1	YV90_MYCPN
96	6	2.6	330	1	GRP2_HUMAN
97	6	2.6	330	1	ODBA_BACSU
98	6	2.6	331	1	GALR_LACCA
99	6	2.6	341	1	RIR2_HELPY
100	6	2.6	342	1	ARGC_STRCO

ALIGNMENTS

RESULT 1

P50031	synecococc
001952	synecocyst
002924	synecocyst
P16571	lymella d
Q94H13	arabidopsis
Q36572	myceteius
Q35873	sturnira li
076537	trichoplusi
P15935	clostridium
Q91109	mus musculu
Q63520	mycobacteri
P77262	escherichia
P31739	aeromonas h
Q98G02	mycoplasma
Q66771	aquifex aeo
Q9TPQ3	thermomaner
Q16994	sus scrofa
Q92na2	delnoccus
Q46607	canis famli
Q75715	homo sapien
P28714	macaca fasc
P42813	arabidopsis
Q73847	prototenus
Q33529	rhizobium l
Q822K5	pyrobaculum
Q96e08	lactococcus
P17596	wolinnella s
Q9paga1	xyella fas
Q9qwm8	mus musculu
P20942	rattus norv
P43584	saccharomyc
Q822Y2	salmonella
Q82kp7	salmonella
P03953	mus musculu
P49275	dermatophag
Q9p103	chlamydia m
P28064	rattus norv
P75266	mycoplasma
P28062	homo sapien
P42369	streptococc
P13309	streptococc
Q99020	mus musculu
P4666	caenorhabdi
Q97766	schizosacch
Q911u8	cavia porce
P6118	azotobacter
P09818	rhizobium m
P06117	bradyrhizob
P00463	bradyrhizob
P33210	saccharomyc
Q91k11	streptomyce
Q91540	foxlow vir
P55172	pseudomonas
Q99433	m putative
Q95859	homo sapien
O66132	buchnera ap
P02649	homo sapien
P10517	macaca fasc
P05770	papio anubi
O89100	m grb2-rela
P22674	homo sapien
P75603	mycoplasma
O75791	h grb2-rela
P37940	bacillus su
O84905	helicobacte
Q92k33	helicobacte
P54855	streptomyce

```
HIS1_LACLA
ID HIS1_LACLA STANDARD; PRT; 208 AA.
AC Q02129; Q9CG94;
DT 01-JUL-1993 (Rel. 26, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DE ATP phosphoribosyltransferase (EC 2.4.2.17).
DE HISG OR L11208
GN Lactococcus lactis (subsp. lactis) (Streptococcus lactis).
OS Bacteria; Firmicutes; Lactobacillales; Streptococcaceae; Lactococcus.
OX NCBI_TaxID=1360;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=NCDO 2118;
RX MEDLINE=93015709; PubMed=1400209;
RA Delorme C., Ehrlich S.D., Renault P.;
RT "Histidine biosynthesis genes in Lactococcus lactis subsp. lactis.";
RL J. Bacteriol. 174:6571-6579(1992).
RN [2]
RP SEQUENCE FROM N.A.
RC STRAIN=IL1403.
RX MEDLINE=21235186; PubMed=11337471;
RA Bolotin A., Winkler P., Mauger S., Jaillon O., Malarne K.,
RA Weissenbach J., Ehrlich S.D., Sorokin A.;
RT "The complete genome sequence of the lactic acid bacterium Lactococcus
RT lactis ssp. lactis IL1403.";
RL Genome Res. 11:731-753(2001).
RN [3]
RP CHARACTERIZATION.
RX MEDLINE=99362697; PubMed=10430882;
RA Sissler M., Delorme C., Bond J., Ehrlich S.D., Renault P.,
RA Fracklyn C.;
RT "An aminoacyl-tRNA synthetase paralog with a catalytic role in
RT histidine biosynthesis.";
RL Proc. Natl. Acad. Sci. U.S.A. 96:8985-8990(1999).
CC -1- CATALYTIC ACTIVITY: L-(5-phospho-D-ribose)-ATP + diphosphate =
CC ATP + 5-phospho-alpha-D-ribose 1-diphosphate.
CC -1- PATHWAY: Histidine biosynthesis; first step. Very important in the
CC regulation of histidine metabolism.
CC -1- SUBUNIT: Homohexamer (By similarity). Binds to hisz possibly to
CC allow the regulation of hisG transferase activity by histidine.
CC -1- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
CC -1- DOMAIN: Lacks the C-terminal regulatory region which is replaced
CC by hisZ.
CC -1- SIMILARITY: BELONGS TO THE ATP PHOSPHORIBOSYLTRANSFERASE FAMILY.
CC SHORT SUBFAMILY.
CC -1- CAUTION: Ref.2 sequence differs from that shown due to a
CC frameshift in position 174.
CC
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation
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CC use by non-profit institutions as long as its content is in no way
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CC or send an email to license@isb-sib.ch).
CC
CC EMBL; U92974; AAB81903.1;
CC EMBL; AE006353; AAK05306.1; ALT_FRAME.
CC PIR; S28532; S28532.
CC PIR; D45734; D45734.
CC InterPro; IPR001348; HisG.
CC Pfam; PF01634; HisG; 1.
CC ProDom; PD003516; HisG; 1.
CC TIGRfams; TIGR00070; hisG; 1.
CC PROSITE; PS01316; ATP_P_PHOSPHORIBOSYLTR; 1.
CC Histidine biosynthesis; Transferase; Glycosyltransferase;
CC COMPLETE PROTEOME.
CC CONFLICT 51 P -> A (IN REF. 1).
CC CONFLICT 86 Y -> D (IN REF. 1).
CC CONFLICT 110 H -> R (IN REF. 1).
CC SEQUENCE 208 AA; 23677 MW; 8CE4CDOA16D39FEF CRC64;

```

```
Query Match 3.0%; Score 7; DB 1; Length 208;
Best Local Similarity 100.0%; Pred. No. 8.8;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 165 DYVELLY 171
DB 80 DYVELLY 86
|||||
|

RESULT 2
HIS5_LEGPN STANDARD; PRT; 213 AA.
AC Q9RDX3;
DT 16-OCT-2001 (Rel. 40, Created)
DT 16-OCT-2001 (Rel. 40, Last sequence update)
DT 15-JUN-2002 (Rel. 41, Last annotation update)
DE Imidazole glycerol phosphate synthase subunit hisH (EC 2.4.2.-) (IGP
DE synthase glutamine amidotransferase subunit) (IGP synthase subunit
DE hisH) (ImGP synthase subunit hisH) (IGPS subunit hisH).
GN HIS5.
OS Legionella pneumophila.
OC Bacteria; Proteobacteria; gamma subdivision; Legionellaceae group;
OC Legionellaceae; Legionella.
OX NCBI_TaxID=446;
RN [1]
RP SEQUENCE FROM N.A.
RC STRAIN=RC1 / Olda / Serogroup 1;
RA Lueneberg E., Zetmann N., Hartmann M., Knirel Y.A., Kooistra O.,
RA Zaehring U., Helbig J., Frosch M.;
RT "A 30 kb gene cluster involved in biosynthesis of the virulence
RT associated lipopolysaccharide carbohydrate moiety of Legionella
RT pneumophila";
RL Submitted (JUN-1998) to the EMBL/GenBank/DBJ databases.
CC -1- FUNCTION: IGPS catalyzes the conversion of PRFAR and glutamine to
CC IGP, AICAR and glutamate. The hisH subunit provides the glutamine
CC amidotransferase activity that produces the ammonia necessary to
CC hisF for the synthesis of IGP and AICAR (By similarity).
CC -1- CATALYTIC ACTIVITY: 5-[(5-phospho-1-deoxyribulose-1-
CC ylaminomethylideneamino)-1-(5-phosphoribosyl)imidazole-4-
CC carboxamide + L-glutamine = imidazole-glycerol phosphate + 5-
CC aminoimidazole-4-carboxamide ribonucleotide + L-glutamate + H(2)O.
CC -1- PATHWAY: Histidine biosynthesis; fifth step.
CC -1- SUBUNIT: Heterodimer of hisH and hisF (By similarity).
CC -1- SUBCELLULAR LOCATION: Cytoplasmic (By similarity).
CC -1- SIMILARITY: CONTAINS 1 TYPE-1 GLUTAMINE AMIDOTRANSFERASE DOMAIN.
CC
CC This SWISS-PROT entry is copyright. It is produced through a collaboration
CC between the Swiss Institute of Bioinformatics and the EMBL outstation
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CC use by non-profit institutions as long as its content is in no way
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CC or send an email to license@isb-sib.ch).
CC
CC EMBL; AJ007311; CAB65214.1;
CC InterPro; IPR000991; GATase_1.
CC Pfam; PF00117; GATase; 1.
CC PROSITE; PS00442; GATASE_TYPE_I; 1.
CC Histidine biosynthesis; Transferase; Glutamine amidotransferase.
CC ACT SITE 81 BY SIMILARITY.
CC ACT SITE 195 BY SIMILARITY.
CC ACT SITE 197 BY SIMILARITY.
CC ACT SITE 197 BY SIMILARITY.
CC SEQUENCE 213 AA; 23310 MW; F27CDEFD7C771D7C CRC64;

```

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Query March 3.0%; Score 7; DB 1; Length 213;
Best Local Similarity 100.0%; Pred. No. 9;
Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 56 NSSVYFV 62
DB 150 NSSVYFV 156
|||||
|

```

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OM protein - protein search, using sw model

Run on: January 13, 2003, 15:44:46 ; Search time 20 Seconds

(without alignments)
1110.353 Million cell updates/sec

Title: US-09-728-911-2

Sequence: 1 MPMKFCFLGFLISFLTGVA.....YQPMIDRRSQRSERCEVIEIP 231

Scoring table: OLIGO
Gapop 60.0 , Gapext 60.0

Searched: 283224 seqs, 96134422 residues

W size : 0

Total number of hits satisfying chosen parameters: 283224

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Listing first 100 summaries

Database : PIR 73:*

1: PIR1:*

2: PIR2:*

3: PIR3:*

4: PIR4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	3.0	180	2	H86775	ATP phosphoribosyl
2	3.0	213	2	B86845	hypothetical prote
3	3.0	218	2	T49885	peptide methionine
4	3.0	236	2	A98351	agropine synthetis
5	3.0	236	2	AE2931	agropine synthetis
6	3.0	241	2	B95856	hypothetical prote
7	3.0	243	2	AC2727	hypothetical prote
8	3.0	243	2	G97508	hypothetical prote
9	3.0	255	2	H72319	mazg protein - The
10	3.0	270	2	JE0274	connexin 31 - huma
11	3.0	303	1	QOEERT	L(+)-tartarate dehy
12	3.0	303	2	H81121	L-tartarate dehydra
13	3.0	303	2	G85966	L-tartarate dehydra
14	3.0	304	2	T32718	hypothetical prote
15	3.0	312	2	T35400	probable phytoene
16	3.0	322	2	AC3207	conserved hypotet
17	3.0	333	2	B90172	conserved hypotet
18	3.0	341	2	DE4565	ribonucleoside dip
19	3.0	357	2	H70346	undecaprenyl-phosp
20	3.0	371	2	T42623	probable sugar tra
21	3.0	446	2	H90094	conserved hypotet
22	3.0	446	2	E96991	hypothetical prote
23	3.0	448	2	E96991	Na+/H+ antiporter,
24	3.0	457	2	S50357	sugar transport pr
25	3.0	479	2	T48025	hypothetical prote
26	3.0	512	2	B69146	glutathione-regula
27	3.0	535	2	S68446	origin recognition
28	3.0	546	2	T40888	probable glucose t
29	3.0	552	2	G96729	hypothetical prote

30	3.0	562	2	E72608	probable huB APE1
31	3.0	567	2	S50708	hexose transport p
32	3.0	567	2	S49600	glucose transport
33	3.0	570	2	S38798	hexose transport p
34	3.0	592	2	S43742	hexose transport p
35	3.0	637	2	C71712	cell division prot
36	3.0	637	2	D97708	cell division prot
37	3.0	788	2	T26967	hypothetical prote
38	3.0	800	2	I51653	deRNA-binding prot
39	3.0	823	2	A93361	endopeptidase Ia (
40	3.0	921	2	G71705	alkaline phosphata
41	3.0	1166	2	S70413	DNA-directed RNA p
42	3.0	1210	2	A48001	phospholipase C (E
43	3.0	2825	2	T14271	Doc4 protein, stre
44	3.0	4273	2	C69679	polyketide synthas
45	2.6	33	2	A60465	cytochrome P450 D1
46	2.6	36	2	F95077	hypothetical prote
47	2.6	56	2	B97945	hypothetical prote
48	2.6	57	2	AH2906	hypothetical prote
49	2.6	82	2	E70972	conserved hypotet
50	2.6	90	2	H95138	probable enoyl-coA
51	2.6	90	2	H95138	hypothetical prote
52	2.6	91	2	H88006	hypothetical prote
53	2.6	91	2	F69252	hypothetical prote
54	2.6	93	2	H81357	hypothetical prote
55	2.6	94	2	AB0949	hypothetical prote
56	2.6	102	2	H87678	hypothetical prote
57	2.6	116	2	G84374	hypothetical prote
58	2.6	118	2	C90583	50S ribosomal prot
59	2.6	121	2	S71007	hypothetical prote
60	2.6	125	2	T16042	hypothetical prote
61	2.6	127	2	PM0464	hypothetical prote
62	2.6	128	2	S73593	hypothetical prote
63	2.6	134	2	B95216	conserved hypotet
64	2.6	136	2	S74683	hypothetical prote
65	2.6	139	2	E40604	hypothetical prote
66	2.6	141	2	S10037	hypothetical prote
67	2.6	141	2	H98079	conserved hypotet
68	2.6	143	2	H81077	hypothetical prote
69	2.6	144	2	S26014	NADH2 dehydrogenas
70	2.6	145	2	S35159	photosystem I chal
71	2.6	148	2	T04727	hypothetical prote
72	2.6	152	2	S36108	hypothetical prote
73	2.6	152	2	G87539	superoxide dismuta
74	2.6	154	2	JC1192	phosphotyrosine pr
75	2.6	154	2	D69019	superoxide dismuta
76	2.6	156	2	D82741	superoxide dismuta
77	2.6	157	2	T15627	conserved hypotet
78	2.6	159	2	AE1241	transcription term
79	2.6	159	2	AE1603	hypothetical prote
80	2.6	160	2	A83110	B. subtilis YqzC p
81	2.6	161	2	B44462	hypothetical prote
82	2.6	161	2	S33624	aliphococyanin be
83	2.6	161	2	E48232	aliphococyanin be
84	2.6	162	2	C31385	cysteine-rich exte
85	2.6	162	2	A81325	allophycocyanin 1
86	2.6	165	2	C48232	probable signal-tr
87	2.6	165	2	G95074	cysteine-rich exte
88	2.6	166	2	F86898	PTS system IIA com
89	2.6	169	2	F72465	single-strand bind
90	2.6	172	2	F83071	hypothetical prote
91	2.6	173	2	S41755	probable transcrip
92	2.6	173	2	B97942	cyclin E type II -
93	2.6	175	2	D86180	hypothetical prote
94	2.6	177	2	T36271	hypothetical prote
95	2.6	178	2	S54444	probable RNA polym
96	2.6	178	2	E88637	preillin 18.1K typ
97	2.6	180	2	PC1305	protein W09G12.6 l
98	2.6	180	2	F71809	genome polypotein
99	2.6	180	2	B64711	hypothetical prote
100	2.6	183	2	E83410	putine nucleoside

ALIGNMENTS

RESULT 1

H86775
 ATP phosphoribosyltransferase (EC 2.4.2.17) [imported] - Lactococcus lactis subsp. lactis
 C:Species: Lactococcus lactis subsp. lactis
 C:Date: 23-Mar-2001 #sequence_revision 23-Mar-2001 #text_change 03-Aug-2001
 C:Accession: H86775
 R:Bolotin, A.; Wincker, P.; Mauger, S.; Jaillon, O.; Malarne, K.; Weissenbach, J.; Ehrlich
 Genome Res. 11, 731-753, 2001
 A:Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis s
 A:Reference number: A86625; MUID:21235186; PMID:11337471
 A:Accession: H86775
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-180 <STO>
 A:Cross-references: GB:AE005176; PID:gl2724177; PIDN:AAK05306.1; GSPDB:GN00146
 A:Experimental source: strain IL1403
 C:Genetics:
 A:Gene: hisG
 C:Keywords: glycosyltransferase; pentosyltransferase

Query Match 3.0%; Score 7; DB 2; Length 180;
 Best Local Similarity 100.0%; Pred. No. 15;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 165 DYVELLY 171
 |||||
 DB 80 DYVELLY 86

RESULT 2

H86845
 hypothetical protein yscB [imported] - Lactococcus lactis subsp. lactis (strain IL1403)
 C:Species: Lactococcus lactis subsp. lactis
 C:Date: 23-Mar-2001 #sequence_revision 23-Mar-2001 #text_change 03-Aug-2001
 C:Accession: B86845
 R:Bolotin, A.; Wincker, P.; Mauger, S.; Jaillon, O.; Malarne, K.; Weissenbach, J.; Ehrlich
 Genome Res. 11, 731-753, 2001
 A:Title: The complete genome sequence of the lactic acid bacterium Lactococcus lactis s
 A:Reference number: A86625; MUID:21235186; PMID:11337471
 A:Accession: B86845
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-213 <STO>
 A:Cross-references: GB:AE005176; PID:gl2724784; PIDN:AAK05860.1; GSPDB:GN00146
 A:Experimental source: strain IL1403
 C:Genetics:
 A:Gene: yscB

Query Match 3.0%; Score 7; DB 2; Length 213;
 Best Local Similarity 100.0%; Pred. No. 17;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 7 FLGLFIS 13
 |||||
 DB 44 FLGLFIS 50

RESULT 3

T49885
 peptide methionine sulfoxide reductase-like protein - Arabidopsis thaliana
 N:Alternate names: protein T211.170
 C:Species: Arabidopsis thaliana (mouse-ear cress)
 C:Date: 02-Jun-2000 #sequence_revision 02-Jun-2000 #text_change 02-Sep-2000
 C:Accession: T49885
 R:Bevan, M.; Murphy, G.; Ridley, P.; Hudson, S.; Bancroft, I.; Mewes, H.W.; Rudd, S.; Le
 submitted to the Protein Sequence Database, April 2000
 A:Reference number: T24493
 A:Accession: T49885
 A:Status: preliminary
 A:Molecule type: DNA

A:Residues: 1-218 <BEV>
 A:Cross-references: EMBL:AL163912; GSPDB:GN00063; ATSP:T211.170
 A:Experimental source: cultivar Columbia; BAC clone T211
 C:Genetics:

A:Gene: ATSP:T211.170
 A:Map position: 5
 A:Introns: 135/3
 C:Superfamily: peptide methionine sulfoxide reductase

Query Match 3.0%; Score 7; DB 2; Length 218;
 Best Local Similarity 100.0%; Pred. No. 18;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 179 SLEKEQK 185
 |||||
 DB 161 SLEKEQK 167

RESULT 4

A98351
 agropine synthesis reductase [imported] - Agrobacterium tumefaciens (strain C58, Dupont)
 C:Species: Agrobacterium tumefaciens
 C:Date: 22-Oct-2001 #sequence_revision 22-Oct-2001 #text_change 11-Jan-2002
 C:Accession: A98351
 R:Goodner, B.; Hinkle, G.; Gattung, S.; Miller, N.; Blanchard, M.; Qurollo, B.; Goldman,
 A.; Liu, F.; Wollam, C.; Allinger, M.; Doughty, D.; Scott, C.; Lappas, C.; Markelz, B.;
 Science 294, 2323-2328, 2001
 A:Title: Genome Sequence of the Plant Pathogen and Biotechnology Agent Agrobacterium tum
 A:Reference number: A97359; PMID:11743194
 A:Accession: A98351
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-236 <KUR>
 A:Cross-references: GB:AE007870; PIDN:AAK90331.1; PID:gl5160366; GSPDB:GN00170
 C:Genetics:
 A:Gene: AGR L 3508
 A:Map position: linear chromosome
 C:Superfamily: ribitol dehydrogenase; short-chain alcohol dehydrogenase homology

Query Match 3.0%; Score 7; DB 2; Length 236;
 Best Local Similarity 100.0%; Pred. No. 19;
 Matches 7; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 106 ASAGSYS 112
 |||||
 DB 147 ASAGSYS 153

RESULT 5

AE2931
 agropine synthesis reductase [imported] - Agrobacterium tumefaciens (strain C58, Dupont)
 C:Species: Agrobacterium tumefaciens
 C:Date: 11-Jan-2002 #sequence_revision 11-Jan-2002 #text_change 01-Feb-2002
 C:Accession: AE2931
 R:Wood, D.W.; Setubal, J.C.; Kaul, R.; Monks, D.; Chen, L.; Wood, G.E.; Chen, Y.; Woo, L
 erage, G.; Gillet, W.; Grant, C.; Guenther, D.; Kutayavin, T.; Levy, R.; Li, M.; McClell
 ; Karp, P.; Romero, P.; Zhang, S.
 Science 294, 2317-2323, 2001
 A:Authors: Yoo, H.; Tao, Y.; Biddle, P.; Jung, M.; Krespan, W.; Perry, M.; Gordon-Kamm, I
 ster, E.W.
 A:Title: The Genome of the Natural Genetic Engineer Agrobacterium tumefaciens C58.
 A:Reference number: AB2577; PMID:11743193
 A:Accession: AE2931
 A:Status: preliminary
 A:Molecule type: DNA
 A:Residues: 1-236 <KUR>
 A:Cross-references: GB:AE008689; PIDN:AAL43867.1; PID:gl7741412; GSPDB:GN00187
 A:Experimental source: strain C58 (Dupont)
 C:Genetics:
 A:Gene: masI
 A:Map position: linear chromosome
 C:Superfamily: ribitol dehydrogenase; short-chain alcohol dehydrogenase homology

QY 106 ASAGSYS 112
 |||||
 DB 147 ASAGSYS 153